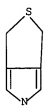
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=> d que 122

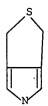
L1 STR



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L5
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L7
             2 SEA FILE=REGISTRY ABB=ON PLU=ON L7 AND L6
L8
             2 SEA FILE=HCAPLUS ABB=ON PLU=ON L8
L10
            32 SEA FILE=HCAPLUS ABB=ON PLU=ON L7
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            32 SEA FILE=HCAPLUS ABB=ON
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L13
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          1265 SEA FILE=HCAPLUS ABB=ON PLU=ON
L16
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                "KATO M TAKAYUKI"/AU OR "KATO MASAHIKO"/AU)
           301 SEA FILE=HCAPLUS ABB=ON PLU=ON
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L17
               AKIRA"/AU)
                                        PLU=ON L16 AND L17
L18
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L19
             4 SEA FILE=HCAPLUS ABB=ON
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                                                (L18 OR L5)
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L20
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L1 STR



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Structure attributes must be viewed using STN Express query preparation.
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L5
              7 SEA FILE=REGISTRY ABB=ON PLU=ON (1313-82-2/BI OR 152940-72-2/
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                63156-10-5/BI OR 646065-36-3/BI)
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L7
             2 SEA FILE=REGISTRY ABB=ON PLU=ON L7 AND L6
L8
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L11
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DUPLICATE IS NOT AVAILABLE IN 'CAOLD'.
ANSWERS FROM THESE FILES WILL BE CONSIDERED UNIQUE
PROCESSING COMPLETED FOR L26
PROCESSING COMPLETED FOR L22
PROCESSING COMPLETED FOR L13
             67 DUP REM L26 L22 L13 (2 DUPLICATES REMOVED)
L32
                ANSWERS '1-2' FROM FILE WPIX
                ANSWERS '3-67' FROM FILE HCAPLUS
=> d all abeq tech 132 1-2;d ibib abs hitstr retable 132 3-67
L32 ANSWER 1 OF 67 WPIX COPYRIGHT 2006 THE THOMSON CORP on STN
     DUPLICATE 1
     2004-122567 [12]
                        WPIX Full-text
AN
DNC
     C2004-049274 [12]
DNN N2004-098082 [12]
     New 3,5-dihydro-1H-thieno(3,4-c)pyrrole derivatives for use in
     e.g. electrical capacitors
     E13; L03; V01
DC
     KANEKO A; KATO M
ΤN
PA
     (NIPS-C) NIPPON SODA CO
CYC 103
     WO 2004005297 A1 20040115 (200412)* JA 21[0]
JP 2004035507 A 20040205 (200412) JA 15
PΙ
                                                          C07D495-04
                                                           C07D495-04
     AU 2003246132 A1 20040123 (200459) EN
                    A1 20050727 (200549) EN
                                                           C07D495-04
     EP 1557419
     US 20050222241 A1 20051006 (200566) EN
                                                           A61K031-407
     WO 2004005297 A1 WO 2003-JP8266 20030630; JP 2004035507 A JP 2002-197401.
     20020705; AU 2003246132 A1 AU 2003-246132 20030630; EP 1557419 A1 EP
     2003-738573 20030630; EP 1557419 A1 WO 2003-JP8266 20030630; US
```

20050222241 A1 WO 2003-JP8266 20030630; US 20050222241 A1 US 2004-520050 20041230 AU 2003246132 Al Based on WO 2004005297 A; EP 1557419 Al Based on WO 2004005297 A PRAI JP 2002-197401 20020705 ICM A61K031-407; C07D495-04 ICS C07D498-02 UPAB: 20060121 WO 2004005297 A1 AΒ NOVELTY - 3,5-Dihydro-1H-thieno(3,4-c)pyrrole derivatives (I) are new. DETAILED DESCRIPTION - 3,5-Dihydro-1H-thieno(3,4-c)pyrrole derivatives of formula (I) are new. R1, R2 = H or 1-10C hydrocarbyl. An INDEPENDENT CLAIM is also included for intermediates of formula (II). Z = organic group. USE - For use in e.g. electrical capacitors. ADVANTAGE - Have good electrical properties e.g. high frequency. CPI: E06-F03; L03-B03A MC EPI: V01-B01B5 TECH ORGANIC CHEMISTRY - Preparation: (I) are prepared e.g. by cyclizing a pyrrole compound of formula (III) to give (II) and deprotecting (II; Z = protecting group). L32 ANSWER 2 OF 67 WPIX COPYRIGHT 2006 THE THOMSON CORP on STN DUPLICATE 2 2004-112968 [12] WPIX Full-text ΑN DNC C2004-046546 [12] DNN N2004-089890 [12] Manufacture of pyrrole derivative having disulfide bond for use as positive electrode active material of secondary battery, involves processing specific compound with base E13: L03: X16 DC KANEKO A; KANEKO T; KATO M IN (NIPS-C) NIPPON SODA CO PA CYC JP 2003335782 A 20031128 (200412)\* JA 6[0] C07D495-04. PΙ ADT JP 2003335782 A JP 2002-138203 20020514 PRAI JP 2002-138203 20020514 IC ICM C07D495-04 ICA C07B061-00 JP 2003335782 A UPAB: 20050528 AB NOVELTY - A compound (II) is processed by a base to obtain a pyrrole derivative (I) having disulfide bond. DETAILED DESCRIPTION - A compound of formula (II) is processed by a base to obtain a pyrrole derivative having disulfide bond of formula (I). R1, R2 = 1-4C linear or branched alkylene; p, q = 0 or 1; andZ1, Z2 = H or organic group both p and q are not 0. An INDEPENDENT CLAIM is included for manufacture of compound of formula (IV), which involves reacting compound of formula (III) with metal sulfide and sulfur in mixture of water and organic solvent under phase-transfer catalysis existence. R1, R2, p, q = as mentioned above; and Z3 = Z1 of formula (I). USE - For manufacturing pyrrole derivative for use as positive

electrode active material of secondary battery.

ADVANTAGE - The *pyrrole* derivative having a disulfide bond is obtained with sufficient yield using an inexpensive reagent. The manufacturing method is industrially advantageous

MC CPI: E06-F03; L03-E01B9

EPI: X16-B01; X16-E01A; X16-E01G

TECH

ORGANIC CHEMISTRY - Preferred Condition: The *pyrrole* derivative is manufactured in inert gas atmosphere.

Preferred Compound: The base is alkali metal hydroxide or alkaline earth metal hydroxide.

L32 ANSWER 3 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:169407 HCAPLUS Full-text

DOCUMENT NUMBER:

142:411126

TITLE:

Sulfolenoporphyrins: synthons for refunctionalization

of porphyrins

AUTHOR(S):

Lee, Sang Hee; Smith, Kevin M.

CORPORATE SOURCE:

Department of Chemistry, Louisiana State University,

Baton Rouge, LA, 70803, USA

SOURCE:

Tetrahedron Letters (2005), 46(12), 2009-2013

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER:

Elsevier B.V.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 142:411126

GI

Using sulfolenopyrroles I (R = H, Rl = CO2CH2Ph) and I (R, Rl = CHO), methods are developed for the synthesis of opp- (e.g II) and adj- (III) bis-sulfolenoporphyrins. Such compds. are useful building blocks for the refunctionalization of the porphyrin system, and readily undergo Diels-Alder cycloaddn. reactions.

IT 144425-36-5 218628-86-5 850424-43-0

850424-47-4

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of sulfolenoporphyrins as synthons for refunctionalization of porphyrins)

RN 144425-36-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 850424-43-0 HCAPLUS

CN 1H-Pyrrole-2-carboxylic acid, 5,5'-(3,5-dihydro-2,2-dioxido-1H-thieno[3,4-c]pyrrole-4,6-diyl)bis[4-ethyl-3-methyl- (9CI) (CA INDEX NAME)

RN 850424-47-4 HCAPLUS

CN 1H-Pyrrole-3-propanoic acid, 2,2'-[(3,5-dihydro-2,2-dioxido-1H-thieno[3,4-c]pyrrole-4,6-diyl)bis(methylene)]bis[5-carboxy-4-methyl-,α,α'-dimethyl ester (9CI) (CA INDEX NAME)

HO2C Me

$$CH_2-CH_2-CH_2$$

OME

 $CH_2$ 
 $CH_2$ 

IT 850424-41-8P 850424-42-9P 850424-44-1P 850424-45-2P 850424-48-5P 850424-49-6P

850424-50-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of sulfolenoporphyrins as synthons for refunctionalization of porphyrins)

RN 850424-41-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4,6-dicarboxaldehyde, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 850424-42-9 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4,6-dimethanamine, 3,5-dihydro-N,N,N',N'-tetramethyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 850424-44-1 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4,6-bis(1,3-benzodithiol-2-yl)-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 850424-45-2 HCAPLUS

CN 1H,3H,23H,25H-Dithieno[3,4-b:3',4'-1]porphine, 7,19-diethyl-12,14-dihydro-8,18-dimethyl-, 2,2,13,13-tetraoxide (9CI) (CA INDEX NAME)

RN 850424-48-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6,6'-(phenylmethylene)bis[3,5-dihydro-, bis(phenylmethyl) ester, 2,2,2',2'-tetraoxide (9CI) (CA INDEX NAME)

RN 850424-49-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6,6'-(phenylmethylene)bis[3,5-dihydro-, 2,2,2',2'-tetraoxide (9CI) (CA INDEX NAME)

RN 850424-50-9 HCAPLUS

CN 23H,25H-Dithieno[3,4-b:3',4'-g]porphine, 10-(3,5-dimethoxyphenyl)1,3,17,19-tetrahydro-21-phenyl-, 2,2,18,18-tetraoxide (9CI) (CA INDEX NAME)

IT 850424-46-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of sulfolenoporphyrins as synthons for refunctionalization of porphyrins)

RN 850424-46-3 HCAPLUS

CN 1H,3H,23H,25H-Dithieno[3,4-b:3',4'-l]porphine-7,19-dipropanoic acid, 12,14-dihydro-8,18-dimethyl-, dimethyl ester, 2,2,13,13-tetraoxide (9CI) (CA INDEX NAME)

RETABLE

Referenced Author | Year | VOL | PG | Referenced Work | Referenced (RAU) | (RPY) | (RVL) | (RPG) | (RWK) | File

=======================================	c+====+====:	=+====	=+=================+===================
Arsenault, G	1960  82	4384	J Am Chem Soc   HCAPLUS
Boudif, A	1996	1235	J Chem Soc, Perkin T   HCAPLUS
Cadamuro, S	1993	12939	J Chem Soc, Perkin T   HCAPLUS
Gunter, M	1999	1803	Chem Commun   HCAPLUS
Gunter, M	12002   6	1673	J Porphyrins Phthalo HCAPLUS
Krautler, B	2000  83	1583	Helv Chim Acta
Kutzki, O	2000  83	12231	Helv Chim Acta   HCAPLUS
Lash, T	12000 12	125	The Porphyrin Handbo HCAPLUS
Montforts, F	12000   39	599	Angew Chem, Int Ed   HCAPLUS
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Nakayama, J	1975	525	J Chem Soc, Perkin T   HCAPLUS
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Nguyen, L	1994  35	7581	Tetrahedron Lett   HCAPLUS
Rieder, A	2000  122	19050	J Am Chem Soc   HCAPLUS
Sessler, J	1987  52	4394	J Org Chem   HCAPLUS
Smith, K	2004  17	1087	Science of Synthesis
Tardieux, C	1998	1267	Synthesis   HCAPLUS
Vicente, M	1997  38	3639	Tetrahedron Lett   HCAPLUS

L32 ANSWER 4 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:146495 HCAPLUS Full-text

DOCUMENT NUMBER:

138:187638

TITLE:

GI

Preparation of (halomethyl) pyrroles from (hydroxymethyl) pyrroles and hydrogen halides

INVENTOR(S): Kato, Masahiko

PATENT ASSIGNEE(S): SOURCE:

Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

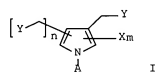
DOCUMENT TYPE:

Patent Japanese

LANGUAGE: 5
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2003055343	A	20030226	JP 2001-250105	20010821
PRIORITY APPLN. INFO.:			JP 2001-250105	20010821
OTHER SOURCE(S):	MARPAT	138:187638		



AB (halomethyl) pyrroles I [A = C1-6 alkyl, (un) substituted Ph, (un) substituted phenylsulfonyl, C1-6 alkylsulfonyl, C1-6 haloalkylsulfonyl, CHO, C2-6 alkylcarbonyl, C1-6 haloalkylcarbonyl, (un) substituted benzoyl; X = halo, C1-6 alkyl, NO2, cyano, (un) substituted phenyl; Y = halo; n = 0, 1; m = 0, 1, 2], useful as intermediates for functional products, agrochems., drugs, etc,m are prepared by treating I (Y = CH2OH; A, X, m, n = same as above) with hydrogen halides. A toluene solution of 1-(p-methylphenylsulfonyl)-3,4-

di(hydroxymethyl) *pyrrole* was treated with HBr solution at room temperature for 5 h to give 90% 1-(p-methylphenylsulfonyl)-3,4-di(bromomethyl) *pyrrole*.

L32 ANSWER 5 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:354172 HCAPLUS Full-text

DOCUMENT NUMBER: 141:106158

TITLE: Theoretical AM1 study of acidity of porphyrins,

azaporphyrins and porphyrazines

AUTHOR(S): Stuzhin, Pavel A.

CORPORATE SOURCE: Department of Organic Chemistry, Ivanovo State

University of Chemical Technology, Ivanovo, 153460,

Russia

SOURCE: Journal of Porphyrins and Phthalocyanines (2003), 7(11

& 12), 813-832

CODEN: JPPHFZ; ISSN: 1088-4246

PUBLISHER: Society of Porphyrins & Phthalocyanines

DOCUMENT TYPE: Journal LANGUAGE: English

The structure-acidity relationship in the series of non-substituted, meso- or AΒ  $\beta$ -substituted and  $\beta$ ,  $\beta$ -annulated porphyrins and porphyrazines have been studied using the AM1 method with UHF basis set. With this purpose, heats of formation have been determined for the geometry optimized structures of the free base macrocycles and corresponding monoanions and dianions formed by deprotonation. Calculated first deprotonation enthalpy values show correlation with available exptl. pKla values and can be used for prediction of acidity. For porphyrazines bearing electron-withdrawing substituents or  $\pi$ -deficient annulated heteroarenes the diamions have lower heats of formation than the corresponding neutral species and such porphyrazines are easily deprotonated upon dissoln. in basic solvents (pyridine, DMF). For porphyrazines with annulated 5-member heteroarenes it is predicted that deprotonation of peripheral NH groups should occur more easily than deprotonation of the internal NH groups. The influence of different types of annulation of 5- and 6-membered heteroarenes to the porphyrazine core on the stability of the macrocyclic system and its acidity are also discussed.

IT 717912-25-9

RL: PRP (Properties)

(dianion, formation enthalpy; theor. AM1 structure-acidity study of porphyrins, azaporphyrins and porphyrazines)

RN 717912-25-9 HCAPLUS

CN 1H,3H,25H,27H-Tetrathieno[3,4-b:3',4'-g:3'',4''-1:3''',4'''-q]porphyrazine-1,3-divl, ion(2-) (9CI) (CA INDEX NAME)

RL: PRP (Properties)

(formation enthalpy and deprotonation enthalpy; theor. AM1 structure-acidity study of porphyrins, azaporphyrins and porphyrazines)

RN 717911-56-3 HCAPLUS

CN 1H,3H,25H,27H-Tetrathieno[3,4-b:3',4'-g:3'',4''-1:3''',4'''-q]porphyrazine-1,3-diyl (9CI) (CA INDEX NAME)

IT 717911-91-6

RL: PRP (Properties)

(monoanion, formation enthalpy; theor. AM1 structure-acidity study of porphyrins, azaporphyrins and porphyrazines)

RN 717911-91-6 HCAPLUS

CN 1H,3H,25H,27H-Tetrathieno[3,4-b:3',4'-g:3'',4''-1:3''',4'''-q]porphyrazine-1,3-diyl, ion(1-) (9CI) (CA INDEX NAME)

RETABLE

Referenced Author	, ,	•	Referenced Work	Referenced
(RAU)		RVL)   (RPG)		File
=======================================	=+=====+==	====+=====	:+=====================================	+========
Anderson, M	1999  38	8  6143	Inorg Chem	HCAPLUS
Andrianov, V	2001  3	107	Advances in Chemistr	•
Arnold, J	12000 13	113	The Porphyrin Handbo	HCAPLUS
Balch, A	1993  32	2  291	Inorg Chem	HCAPLUS
Berezin, B	1977  20	0  807	Izv Vyssh Uchebn Zav	HCAPLUS
Bernauer, K	1962  4	5  2487	Helv Chim Acta	HCAPLUS
Bilton, J	1937	1922	J Chem Soc	HCAPLUS
Bordwell, F	1988  23	1  456	Acc Chem Res	HCAPLUS
Brigaud, O	1992  1	6  1031	New J Chem	.HCAPLUS
Burk, P	1993  8	6  417	Theoret Chim Acta	HCAPLUS

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                                           |Thesis; Ivanovo Inst|
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Donzello, M
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Fitzgerald, J
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Gal'Pern, M
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Ghosh, A
                       12000 17
                                           |The Porphyrin Handbo|HCAPLUS
Ghosh, A
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Kobayashi, N
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McEwen, W
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Morley, J
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Petrov, O
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Richards, R
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Sheinin, V
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Silvers, S
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                                           |Coord Chem Rev
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Stuzhin, P
                                    1500
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Zandler, M
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L32 ANSWER 6 OF 67 HCAPLUS COPYRIGHT 2006 ACS. on STN
                        2002:975665 HCAPLUS Full-text
ACCESSION NUMBER:
                        138:24715
DOCUMENT NUMBER:
TITLE:
```

Kaneko, Akira; Kato, Masahiko; INVENTOR(S):

Tsubokura, Shiro

PATENT ASSIGNEE(S):

Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

LANGUAGE:

SOURCE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2002371068	Α	20021226	JP 2001-184932	20010619
PRIORITY APPLN. INFO.:			JP 2001-184932	20010619

OTHER SOURCE(S):

CASREACT 138:24715; MARPAT 138:24715

GΙ

AB Title compds. I (A = H, hydrocarbyl; B = H, alkyl) are prepared by decarboxylation of cyanoimidazolecarboxylic acids II in organic solvents in the presence of phosphonium or ammonium salt catalysts and alkali or alkaline earth metal salts. Thus, heating 4(5)-cyano-5(4)-imidazolecarboxylic acid in o-dichlorobenzene in the presence of tetrabutylphosphonium bromide and LiCl gave 89% 4(5)-cyanoimidazole.

L32 ANSWER 7 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:847766 HCAPLUS Full-text

DOCUMENT NUMBER:

137:326819

TITLE:

Process for manufacturing 4(5)-cyanoimidazole

derivatives

INVENTOR(S):

Kato, Masahiko; Tsubokura, Shiro;

Kaneko, Akira

PATENT ASSIGNEE(S):

Nippon Soda Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2002322158	A	20021108	JP 2001-128653	20010426
PRIORITY APPLN. INFO.:		- 107 206010	JP 2001-128.653	20010426

OTHER SOURCE(S): MARPAT 137:326819

GΙ

$$Y-N$$
 $X$ 
 $X$ 
 $I$ 

The title compds. I [R = H; X = H, alkyl; Y = H, alkyl, etc.], useful as AB intermediates for pharmaceuticals and agrochems., are prepared by heating I [R = CO2H; X, Y = as defined above] in an organic solvent with gradual temperature increase (in the range of 160° to 220°). Thus, a mixture of 4cyanoimidazole-5-carboxylic acid in nitrobenzene was heated to 160° over 2.5 h; the reaction mixture was then heated from  $160^{\circ}$  to  $165^{\circ}$  over 0.5 h; the reaction mixture was then heated to 170°; the reaction mixture was heated at 170° for 2 h and then heated to 175°; the reaction mixture was heated at 175° for 2 h; the reaction mixture was then heated to 180° and kept at 180° for 2 h to give, after workup, 4-cyanoimidazole in 53% yield. The title process is safe.

L32 ANSWER 8 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:384316 HCAPLUS Full-text

DOCUMENT NUMBER:

136:386121

TITLE:

Method for preparation of disulfide-containing

pyrroles by oxidative cyclization of

3,4-bis(mercaptoalkyl)pyrroles

INVENTOR(S):

Kato, Masahiko; Iihama, Teruyuki

PATENT ASSIGNEE(S):

Nippon Soda Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF Patent

DOCUMENT TYPE:

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2002145884 PRIORITY APPLN. INFO.:	A	20020522	JP 2000-335992 JP 2000-335992	20001102 20001102
OTHER SOURCE(S):	CASREA	ACT 136:38612	1; MARPAT 136:386121	
GI				

Pyrrole derivs. containing an intramol. disulfide bond (I; R1, R2 = linear or AΒ branched C1-4 alkylene) are prepared by air oxidation of 3,4bis(mercaptoalkyl)pyrrole derivs. (II; R1, R2 = same as above; Z = protecting group) under basic conditions, optionally in the presence of at least one catalyst selected from iron(III) or copper(II) salt catalyst, followed by deprotection. This process gives I in a simple procedure without using expensive reagents. I are monomers for conductive polypyrroles which are useful as cathode material for high-capacity secondary battery. Thus, H2S was blown into a solution of 1.23 g NaOEt in 20 mL ethanol under cooling at 5-0  $^{\circ}$ to prepare an ethanol solution of Na2S which was treated with 0.3 g 1-tertbutoxycarbonyl-3,4- bis(chloromethyl)pyrrole and allowed to react at room temperature for 4 h while blowing H2S into the reaction mixture To reaction mixture were added a solution of 0.11 g NaOEt in ethanol and then a catalytic amount of FeCl3 and the resulting mixture was allowed to react at room temperature for 30 min while blowing air into the reaction mixture and then at room temperature for 22 h after stopping blowing air to give 88% 6-tertbutoxycarbonyl-4,6- dihydro-1H-dithiino[4,5-c]pyrrole which was stirred with NaOMe at room temperature for 22 h to give 28% 4,6-dihydro-1H-dithiino[4,5-c] pyrrole, i.e. I (R1 = R2 = CH2).

L32 ANSWER 9 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:359892 HCAPLUS Full-text

DOCUMENT NUMBER:

136:378253

TITLE:

Polysulfide-containing polypyrrole and

manufacture of the polymer

INVENTOR(S):

Koyama, Noboru; Kato, Masahiko; Kaneko,

Takehiko

PATENT ASSIGNEE(S):

SOURCE:

Nippon Soda Co., Ltd., Japan

Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

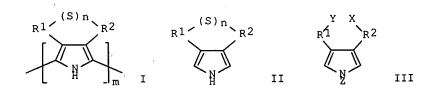
LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE		APPLICATION NO.	DATE	
	<u>-</u> -				
JP 2002138134 PRIORITY APPLN. INFO.: OTHER SOURCE(S):	А маррат	20020514 136:378253	JP 2000-335991 JP 2000-335991	20001102 20001102	
GI	11111(1111	100.070200			



The elec. conductive polysulfide-containing polypyrrole is that represented as I (R1, R2 = linear or branched C1-4 alkylene; n = 3-6; m = 10-10,000). The polymer is manufactured by electrolytic polymerization of a pyrrole II (R1, R2, and n are the same as I), which is manufactured by reaction of another pyrrole III (R1 and R2 are the same as I; X, Y = halogen; Z = protecting

group), alkali metal sulfide, and S. The polymer is regarded as a candidate for cathode active mass in secondary batteries.

L32 ANSWER 10 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:151529 HCAPLUS Full-text

DOCUMENT NUMBER: 139:6702

TITLE: Establishing a library of porphyrin building blocks

for superstructured assemblies: Porphyrin dienes and

dienophiles for cycloaddition reactions

AUTHOR(S): Gunter, Maxwell J.; Tang, Hesheng; Warrener, Ronald N.

CORPORATE SOURCE: Division of Chemistry, University of New England,

Armidale, NSW 2351, Australia

SOURCE: Journal of Porphyrins and Phthalocyanines (2002), 6(11

& 12), 673-684

CODEN: JPPHFZ; ISSN: 1088-4246

PUBLISHER: Society of Porphyrins & Phthalocyanines

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:6702

The synthesis and utility of a series of porphyrins with (masked) diene and dienophile functionality are described. The key porphyrin diene is synthesized from a sulfolenopyrrole by a 3+1 strategy. A range of Diels-Alder cycloadducts is readily accessed from the diene by mild thermal extrusion of sulfur dioxide from the sulfolenoporphyrin, which produces the reactive porphodimethylidene. Each of these cycloadducts is fused to the porphyrin nucleus through a cyclohexene ring thus retaining some conformational flexibility in the resultant structures. The structures can be rigidified by mild oxidation to the corresponding benzo-derivs. Diels-Alder reaction of the porphyrin 1,3-diene resulting from the sulfolenoporphyrin with norbornadiene produces the norbornene derivative, which can serve as a dienophile or dipolarophile in subsequent cycloaddn. reactions. Nevertheless, a preferred route to this structure is through a corresponding 1+3 route, where the norbornene component is part of the tripyrrane. Extension of the synthetic protocols allows ready access to a "mixed function" porphyrin, containing both diene and dienophile components. Likewise, the synthesis of a bis-norbornene porphyrin is described. A collection of each of these reactive components is the basis for a library of building blocks which allows easy and simple entry to a wide variety of complex porphyrin-containing superstructures.

IT 190449-12-8

RL: RCT (Reactant); RACT (Reactant or reagent)
(establishing a library of porphyrin dienes and dienophiles for cycloaddn. reactions)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

234437-54-8P 262611-68-7P 267237-68-3P 532994-06-2P 532994-08-4P 532994-10-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(establishing a library of porphyrin dienes and dienophiles for cycloaddn. reactions)

RN 144425-36-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 234096-96-9 HCAPLUS

CN 1H-Pyrrole-2-carboxylic acid, 5,5'-[(3,5-dihydro-2,2-dioxido-1H-thieno[3,4-c]pyrrole-4,6-diyl)bis(methylene)]bis[4-ethyl-3-methyl-, bis(phenylmethyl) ester (9CI) (CA INDEX NAME)

RN 234437-54-8 HCAPLUS

CN 22H,24H-Thieno[3,4-b]porphine, 7,18-diethyl-1,3-dihydro-8,17-dimethyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 262611-68-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6-formyl-3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 267237-68-3 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 532994-06-2 HCAPLUS

CN 13,16-Methano-26H,28H-naphtho[2,3-b]thieno[3,4-l]porphine, 7,22-diethyl-1,3,12,12a,13,16,16a,17-octahydro-8,21-dimethyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 532994-08-4 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-6-(hydroxymethyl)-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 532994-10-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-4,6-diiodo-, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 532994-09-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (establishing a library of porphyrin dienes and dienophiles for cycloaddn. reactions)

RN 532994-09-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6,6'-[(4,5,6,7-tetrahydro-5,8-methano-2H-isoindole-1,3-diyl)bis(methylene)]bis[3,5-dihydro-,bis(phenylmethyl) ester, 2,2,2',2'-tetraoxide (9CI) (CA INDEX NAME)

RETABLE					
· Referenced Author	Year	VOL	PG	Referenced Work	
	(RPY)			(RWK)	File
				+=========	
·	12000	•	•		HCAPLUS
	1996		-	Comprehensive Supram	
Bell, T	12000	122	•	1	HCAPLUS
Butler, D	1998	l		Synlett	
Clyde-Watson, Z	1998	122	1135	New J Chem	HCAPLUS
Crossley, M	2002	1	1122	Chem Commun (Cambrid	HCAPLUS
Crossley, M	1996	1	12675	Chem Soc, Perkin Tra	HCAPLUS
	1991		1569	J Chem Soc, Chem Com	HCAPLUS
Crossley, M	11995	1	2379	J Chem Soc, Chem Com	HCAPLUS
	1996	37	3569	Tetrahedron Lett	HCAPLUS
	2001	25	1368	New J Chem	HCAPLUS
	1999	38	3219	Angew Chem, Int Ed	HCAPLUS
	11998	1	11661	Chem Commun (Cambrid	HCAPLUS
	2000	I	893	Chem Commun (Cambrid	HCAPLUS
	11997	İ	3161	J Chem Soc, Perkin T	HCAPLUS
•		142	45	Tetrahedron Lett	HCAPLUS
	1998	1	2739	Chem Commun (Cambrid	HCAPLUS
	2001	16	406	Molecules	HCAPLUS
	2002	4	12165	Org Lett	HCAPLUS
Johnston, M	2002	58	3445	Tetrahedron	HCAPLUS
	1998	37	916	Angew Chem, Int Ed	HCAPLUS
	11996	37	5931	Tetrahedron Lett	HCAPLUS
The state of the s	1998	37	2368	Angew Chem, Int Ed	HCAPLUS
	1994	35	1995	Tetrahedron Lett	HCAPLUS
	2000	1122	5220	J Am Chem Soc .	HCAPLUS
	1997	36	361	Angew Chem, Int Ed	HCAPLUS
	1998	l	11	Chem Commun (Cambrid	HCAPLUS
	11999	64	6653	J Org Chem	HCAPLUS
Sanders, J	11998	4	11378	ChemEur J	HCAPLUS
Silva, A	1999	1	1767	Chem Commun (Cambrid	HCAPLUS
	2002	167	1726	J Org Chem	HCAPLUS
Silva, A	2000	141	13065	Tetrahedron Lett	HCAPLUS
	11997	1	1199	Chem Commun (Cambrid	HCAPLUS
Vicente, M	1997	38	3639 ·	Tetrahedron Lett	HCAPLUS
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the state of the s	1998	I	585	Synlett	HCAPLUS
•					

Warrener, R | 1998 | | 593 | Synlett | HCAPLUS | 2heng, G | 11996 | | 1119 | Chem Lett | HCAPLUS |

L32 ANSWER 11 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2003:151528 HCAPLUS Full-text

DOCUMENT NUMBER: 139:6701

TITLE: Porphyrin building blocks: Using a modified

Barton-Zard approach to construct annulated pyrroles Johnstone, Ken D.; Pearce, Wayne A.; Pyke, Simon M. Department of Chemistry, The University of Adelaide,

CORPORATE SOURCE: Department of Chem: SA 5005, Australia

SOURCE: Journal of Porphyrins and Phthalocyanines (2002), 6(11

& 12), 661-672

CODEN: JPPHFZ; ISSN: 1088-4246

PUBLISHER: Society of Porphyrins & Phthalocyanines

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 139:6701

AB A modification of the Barton-Zard pyrrole synthesis involving condensation of isocyanoacetate esters with cyclic unsatd. sulfones using sodium hydride as base is demonstrated for the construction of annulated pyrroles.

IT 190449-12-8P

AUTHOR(S):

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of porphyrin building blocks using a modified Barton-Zard approach to construct annulated pyrroles)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE			
Referenced Author	Year   VOL   PG	Referenced Work	Referenced
(RAU)	(RPY) (RVL) (RPG	G)   (RWK)	File
=======================================	=+====+====	:==+===================================	=+=======
Abel, Y	1998  81  1978	Helv Chim Acta	HCAPLUS
Arnold, D	1994  47  969	Aust J Chem	HCAPLUS
Arnold, D	1999  52  421	Aust J Chem	HCAPLUS
Bag, N	1995  36  6409	Tetrahedron Lett	HCAPLUS
Bailey, W	1954   76   1932	!  J Am Chem Soc	HCAPLUS
Barton, D	1985    1098	J Chem Soc, Chem Co	m HCAPLUS
Barton, D	1990  46  7587	Tetrahedron	HCAPLUS
Bitha, P	1988  25  1035	J Heterocycl Chem	HCAPLUS
Bobal, P	2001  38  527	J Heterocycl Chem	HCAPLUS
Boelle, J	1997    1451	.  Synthesis	HCAPLUS
Burns, D	1995  25  379	Synth Com	HCAPLUS
Chayer, S	2001   42   17759	Tetrahedron Lett	HCAPLUS
Chen, Q	2002  32  1031	.  Synth Com	HCAPLUS
Chen, S	1997  34  273	J Heterocycl Chem	HCAPLUS
Cheng, W	2001   66   5528	3  J Org Chem	HCAPLUS

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Inomata, K
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Ito, S
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Ito, S
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Kwart, H
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Lash, T
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Lash, T
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Lash, T
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May, D
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Meinwald, J
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Mueller, W
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Ono, N
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Ono, N
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Padwa, A
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Padwa, A
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Palmer, B
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Pearce, W
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Pelkey, E
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Pelkey, E
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Perrin, D
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Tang, J
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Uno, H
                                  |4347 | J Chem Soc, Perkin T|HCAPLUS
Uno, H
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                                                            HCAPLUS
                                  |3639 |Tetrahedron Lett
                                                             | HCAPLUS
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Vicente, M
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L32 ANSWER 12 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2002:881502 HCAPLUS Full-text DOCUMENT NUMBER: 139:230515
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TITLE: Product class 13: 1H-pyrroles

AUTHOR(S): Black, D. StC.

CORPORATE SOURCE: School of Chemistry, The University of New South

Wales, Sydney, 2052, Australia

SOURCE: Science of Synthesis (2002), 9, 441-552

CODEN: SSCYJ9

PUBLISHER: Georg Thieme Verlag
DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review describing the preparation of 1H-pyrroles. Covered reactions include ring transformation and ring-closure reactions, substituent modifications.

IT 190449-12-8P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of pyrroles via ring-closure reactions, ring transformations, and substituent modifications)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

DETABLE

RETABLE					
	Year			Referenced Work	Referenced
	(RPY)			(RWK)	File
=======================================					+=======
	•		11329		HCAPLUS
Adamczyk, M	1996	52		·	HCAPLUS
Adler, A	1967	32		•	HCAPLUS
Alazard, J	1993	130	779	Bull Soc Chim Fr	HCAPLUS
Alberola, A	1990	31			HCAPLUS
Alberola, A	1990		2681	J Chem Soc, Perkin T	HCAPLUS
Aldabbagh, F	1997	38	7937	Tetrahedron Lett	HCAPLUS
Almerico, A	1994	37	1549	Heterocycles	HCAPLUS
Alper, H	11977	99	14330	J Am Chem Soc	HCAPLUS
Alvarez, A	11992	57	1653	J Org Chem	HCAPLUS
Amarnath, V	1991	56	6924	J Org Chem	HCAPLUS
Andersen, K	1993	136	1716	J Med Chem	HCAPLUS
Anderson, H	1965	43	409	Can J Chem	HCAPLUS
Anderson, H	11967	45	897	Can J Chem	HCAPLUS
Anderson, W	11986	16	387	Synth Commun	l <u>.</u>
Angelini, G	11975	105	961	Gazz Chim Ital	HCAPLUS
Angelini, G	11980	102	1377	J Am Chem Soc	HCAPLUS
Angelini, G	1980	45	1786	J Org Chem .	HCAPLUS
Anon	1950	16		Heterocyclic Compoun	
Anon	1990	1		The Chemistry of Het	
Antonio, Y	1994	172	15	Can J Chem	HCAPLUS
Aoyagi, K	1988	İ	1891	Chem Lett	HCAPLUS
Aoyagi, Y	1996	137	19203	,	HCAPLUS
Apparao, S	1981		65		HCAPLUS
Arcadi, A	1997	1	667		HCAPLUS
Artis, D	1992	170	1838	•	HCAPLUS
Artis, D	1994	59	2456	J Org Chem	HCAPLUS
Attanasi, O	1993	••	315	J Chem Soc, Perkin T	
Attanasi, O	1986	•	1299	Org Prep Proced Int	HCAPLUS
Attanasi, O	1997	1	1128.	1 - 2	HCAPLUS
Aubert, T	1989	1	1369	J Chem Soc, Perkin T	HCAPLUS
Avalos, M	1989	•	C7		HCAPLUS
Baccolini, G	1987		788	J Chem Soc, Chem Com	
			6817		HCAPLUS
Baciocchi, E	11993		7610		HCAPLUS
Baciocchi, E	11993	•	13799	•	HCAPLUS
Backvall, J	1981	•	59 ·	J Chem Soc, Chem Com	•
Badger, G	1964	117	1987	Aust J Chem	HCAPLUS

5	11002	1	1604	J Chem Soc, Chem Com   H	CADIUS
Baldwin, J	11982		624	•	ICAPLUS
Baltazzi, E	1963		511	,	ICAPLUS
Barcock, R	•	•	11187	•	CAPLUS
Barker, P	-		14849	J Org Chem	CAPTIIC
Barluenga, J	•	•	11696		ICAPLUS
Barluenga, J	•	161	12185		ICAPLUS
Barluenga, J		62	19229		ICAPLUS
Barluenga, J	11975	1	1642	•	ICAPLUS.
Barnett, G		58	1409	·	ICAPLUS
Barton, D	•	1	1098	J Chem Soc, Chem Com   H	
Barton, D	1986	•	12243	J Chem Soc, Perkin T   H	
Barton, D	1990	146 .	7587	•	ICAPLUS
Battersby, A	11983		11240	J Chem Soc, Chem Com   H	
Bauer, H	1970	736	11	Justus Liebigs Ann C H	
Baumes, R	1974		1147		ICAPLUS
Baxter, A	11994		207	Synthesis   H	ICAPLUS
Bayer, H	1970	103	2356	•	ICAPLUS .
Bean, G	11967	32	228	•	ICAPLUS
Bellamy, F	1975	3	395	Heterocycles  H	ICAPLUS
Benages, I	1978	43	4273	J Org Chem  H	ICAPLUS
Bertschy, H	1990	102	1828	Angew Chem  H	ICAPLUS
Bertschy, H	1990	129	1777	Angew Chem Int Ed En	
Beveridge, S	1971	124	1229	Aust J Chem   H	HCAPLUS
Black, D		42	71	Aust J Chem   H	ICAPLUS
Black, D	11997	153	8565	Tetrahedron  H	CAPLUS
Black, D	-	153	18573	Tetrahedron   H	CAPLUS
Boberg, F	1984	i	911	Liebigs Ann Chem  H	<b>ICAPLUS</b>
Boberg, F	11985	i ·	1239		<b>ICAPLUS</b>
Boelle, J	11997	i	1451	-	HCAPLUS
Bogdal, D	11997	•	71.5	<del>-</del>	HCAPLUS
Bohm, S	1989		200	Collect Czech Chem C H	<b>ICAPLUS</b>
Bonnett, G	1963	İ		J Chem Soc	
Bordner, J	11965	30	3824	J Org Chem	
Boukou-Poba, J	11979	i	1717	· •	CAPLUS
Bray, B		55	6317	·	HCAPLUS
Brennan, M	11986	24	2879	•	HCAPLUS
Brimble, M	11990	1	311	J Chem Soc, Perkin T   H	
Broadbent, H		5	757	· ·	HCAPLUS
Bruekelman, S	11984	1	2801	J Chem Soc, Perkin T H	
Buchwald, S	11989	•	14486		HCAPLUS
Buchwald, S.	11989		1776	·	HCAPLUS
Bullock, E	11958		11430	•	HCAPLUS
Bumagin, N	11995	•	11537	Zh Org Khim	
		119	1345	-	HCAPLUS
Bundgaard, T		125	1379		HCAPLUS
Burns, D	11993	123	273	J Chem Soc, Perkin T F	
Cadamuro, S	11993	1	12939	J Chem Soc, Perkin T F	
Cadamuro, S			1805		HCAPLUS
Caddick, S	11992		•	<del>-</del>	HCAPLUS
Cambie, R	1990		11923		HCAPLUS
Campi, E	-	45	1167  2563	J Chem Soc C	ICAL DOD
Candy, C	1970	112		Organometallics	
Carrd, F		112	2478  2706	J Am Chem Soc	
Carter, P	•	109			HCAPLUS
Cartoon, M	•	234	123  155	Comprehensive Hetero	TOWERDS
Chadwick, D	•	4	11343	J Chem Soc, Perkin T	HCADTITE
Chadwick, D	11982			J Chem Soc, Perkin T	
Chadwick, D	1983	_	193		
Chahma, M	11994		1366	•	HCAPLUS
Chakrabarti, J	•	11	417		HCAPLUS
Chamberlin, K	11979	ITZ	1567	Heterocycles	HCAPLUS

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Chamberlin, K	1978	8	579	Synth Commun	HCAPLUS
Chan, H	1997	1	1515	Chem Commun (Cambrid	HCAPLUS
, · ·	1994		•	J Chem Soc, Perkin T	
					HCAPLUS
Chapelle, J	1971		1280	Bull Soc Chim Fr	HCAPLUS
Chatani, N	1986	27 ·	4201	Tetrahedron Lett	HCAPLUS
			•		HCAPLUS
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· · · · - / ~				•	HCAPLUS
Chen, W	1992	70	151	Org Synth	HCAPLUS
Cheng, D	1976	13	1145	J Heterocycl Chem	HCAPLUS
	11976	13		_	<b>HCAPLUS</b>
	11977			· -	HCAPLUS
	•				
Childs, R	1966		•		HCAPLUS
Chiu, P	1988	44	3531	Tetrahedron	HCAPLUS
Chiusoli, G	1989		262	Synthesis	HCAPLUS
•				<del></del>	HCAPLUS
				-	
	•		•	•	HCAPLUS
Chou, S	1991		-	• •	HCAPLUS
Cirrincione, G	1997		1169	Synthesis	HCAPLUS
	1982	147	5217	J Org Chem	HCAPLUS
		•	566	•	HCAPLUS
		•			
· · · - <b>3 ·</b>		•	•	·	HCAPLUS
Cooney, J	1981	46	2570		HCAPLUS
Cooney, J	1983	15	292	Org Prep Proced Int	HCAPLUS
<b>2</b> •			7159	Tetrahedron Lett	HCAPLUS
• •		-	4603	•	HCAPLUS
· · ·	-			•	
· · · · ·	•	•	11974	•	
Cristau, H	1985			•	HCAPLUS
Crump, D	11977	142	105	J Org Chem	HCAPLUS
	-	24	1793	J Heterocycl Chem	HCAPLUS
	1990	1	761	J Chem Soc, Perkin T	
		1 0 0			1
Danks, T	-	29	11425	Tetrahedron Lett	
Darbeau, R	1997	162	18091	· · · · · ·	HCAPLUS
Dave, C	1987	64	713	J Indian Chem Soc	HCAPLUS
Dave, C	11987	64	713	J Indian Chem Soc	HCAPLUS
Davidson, D			361	•	HCAPLUS
			15696		HCAPLUS
Davies, H					•
Davies, H	•		5203		HCAPLUS
Davies, H	1989	30	4653	Tetrahedron Lett	HCAPLUS
DeShong, P	1985	50	12309	J Org Chem	HCAPLUS
De Kimpe, N	11984	i I	182		HCAPLUS
		61	18730	•	HCAPLUS
De Leon, C					1
De Leon, C	•	13	7731	Tetrahedron	! ! a = = = :: a
De Rosa, M		l	1757	J Chem Soc, Chem Com	HCAPLUS
De Rosa, M	1989	4	5347	J Org Chem	
De Rosa, M		136	19261	Tetrahedron Lett	HCAPLUS
	1997	1	825	·	HCAPLUS
Dekura, F		133		•	•
Del Valle, J	•	32	1899	<u>-</u>	HCAPLUS
Dell'Erba, C	1995	51	5181		HCAPLUS
Demopoulos, V	1988	125	635		HCAPLUS
Demopoulos, V	1986	118	1278	Org Prep Proced Int	HCAPLUS
Dhanak, D	11986	i	2181	J Chem Soc, Perkin T	
		126	11839		HCAPLUS
Di Santo, R					
Dickinson, C		27	12470		HCAPLUS
Dimroth, K	11961	639	102	Justus Liebigs Ann C	HCAPLUS
Dischler, B	1961	16	1180	Z Naturforsch, Teil	1
Dragisich, V		1112	1251	J Am Chem Soc	HCAPLUS
=		31	1255	J Heterocycl Chem	HCAPLUS
Drinan, M		1 2 7			
Duhamel, L	11974	l	3167	Tetrahedron Lett	HCAPLUS
Duhamel, P	1973	I	1339	Tetrahedron Lett	HCAPLUS

	. 1 0 0 5			17.01	
Dumoulin, H	1995		1703	·	HCAPLUS
Dumoulin, H	1997	34	13	J Heterocycl Chem	HCAPLUS
D'Auria, M	1997	1	2369	J Chem Soc, Perkin T	HCAPLUS
Earle, M	1990	i 31	14229	•	HCAPLUS
Eberlin, M		153	2084	•	HCAPLUS
•	11984	-		<del>-</del>	
Edwards, M		49	13503		HCAPLUS
Elguero, J	1974	139	357	_	HGAPLUS
Elming, N	1952	6	1867	Acta Chem Scand	HCAPLUS
Engel, N	1978	190	719	Angew Chem	HCAPLUS
Engel, N	11978	17	676	Angew Chem Int Ed En	
Eyley, S	1988	29	2997	_	HCAPLUS
Fabiano, E	1991	1	3371	J Chem Soc, Perkin T	
		•			
Fan, J	1997	16	4232		HCAPLUS
Fang, Y		125	1857		HCAPLUS
Fanghanel, E	1969	311	388	J Prakt Chem	
Fegley, M	1957	179	4144	J Am Chem Soc	HCAPLUS
Fischer, H	1934	13	1	Die Chemie des Pyrro	HCAPLUS
Fischer, H		II	1202	Org Synth, Coll	
Fischer, H		III	1217	Org Synth, Coll	
					UCARTUC
Friedman, M	•	130	1859	_	HCAPLUS
Fritz, H	1971	744	81	Justus Liebigs Ann C	
Fuhrhop, J	1985		1699	Liebigs Ann Chem	
Fujii, H	1997	38	1427	Tetrahedron Lett	HCAPLUS
Fursmer, A	1995	160	16637	J Org Chem	
Furstner, A		13	11047	Tetrahedron Lett	
Furusho, Y	1996	1	1183	J Chem Soc, Perkin T	HCAPLUS
•		130	17007		HCAPLUS
Gage, J		138			
Ganske, J		54	4801		HCAPLUS
Gao, Y		37	17787		HCAPLUS
Gewald, K	1976.	318	1663	J Prakt Chem	HCAPLUS
Gewald, K	1992	1334	1491	J Prakt Chem	HCAPLUS
Ghosez, L	11977	17	1895	Heterocycles	HCAPLUS
Gilchrist, T	1997	ĺ	13005	J Chem Soc, Perkin T	HCAPLUS
Gilow, H	-	28	1025		HCAPLUS
	•		12221	<del>-</del>	HCAPLUS
Gilow, H	1981	146			
Girard, Y		148	13220	. 2	HCAPLUS
Gjos, N		125	12596	•	
Gompper, R	11979	1	213	• •	HCAPLUS
Gossauer, A	11974	1	1	Die Chemie der Pyrro	
Gossauer, A	11976	159	1698	Helv Chim Acta	HCAPLUS
Gotthardt, H		103	12625		HCAPLUS
Grandberg, I		15	501	Chem Heterocycl Comp	
Grandberg, I		115	620	Khim Geterotsikl Soe	
				·	
Greenhouse, R		150	12961	•	HCAPLUS
Grigg, R	1990	46	4003		HCAPLUS
Grob, C	1953	136	49		HCAPLUS
Gronowitz, S	1961	126	12615	J Org Chem	HCAPLUS
Gupton, J	1990	55	14735	J Org Chem	HCAPLUS
Gupton, J	1992	57	15480	J Org Chem	HCAPLUS
Gupton, J		52	6879	· ·	HCAPLUS
Gupton, J	-	54	5075		HCAPLUS
•		35	1843		HCAPLUS
Hamby, J				<del>-</del>	
Hamdan, A		13	741	• -	HCAPLUS
Hantzsch, A		123	1474	Ber Dtsch Chem Ges	1
Harbuck, J	1971	36	853	J Org Chem	l
Harris, R	1972	125	1985	Aust J Chem	HCAPLUS
Harsanyi, M	11987	152	12209		HCAPLUS
Hasan, I		146	157	<del>-</del>	HCAPLUS
Hauptmann, S		314	353	-	HCAPLUS
Hauptmann, S	11968	1	1333		HCAPLUS
naupemann, 5	11300	1 .	LICTI	Liectanearon Deff	HOMPHOS

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	•			Z Chem	HCAPLUS
Hauptmann, S	1969	19	122	Z Chem	
Heaney, H	1988	1	1161	J Chem Soc, Chem Com	HCAPLUS
Heaney, H	1973		499	J Chem Soc, Perkin T	HCAPLUS
	1966	31	3924	J Org Chem	HCAPLUS
· - · · •	1977				HCAPLUS
· - • • • • • • • • • • • • • • • • • •		186	107	J Am Chem Soc	HCAPLUS
				•	HCAPLUS
	1988			• • • •	HCAPLUS
•				-	HCAPLUS
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3.	•	-		. •	HCAPLUS
					HCAPLUS
	•		•	J Am Chem Soc	
Hollins, R	•	-		•	HCAPLUS
Holy, A	1965	130	1346	Collect Czech Chem C	HCAPLUS
Hombrecher, H	1990	1	1062	Synthesis .	
Hou, D	1992	75	12608	Helv Chim Acta	HCAPLUS
	1981	18	1127	J Heterocycl Chem	HCAPLUS
3.			1133	J Heterocycl Chem	HCAPLUS
3.	1976				HCAPLUS
3.	11996		1893	J Chem Soc, Perkin T	•
	11994	•	51		HCAPLUS
	•	•	2611	• •	HCAPLUS
	-	•		•	HCAPLUS
•	•	•			
	•	-	659	Chem Heterocycl Comp	
,		•	1790	Khim Geterotsikl Soe	
•	,	1	1737		HCAPLUS
1 ,			2577	•	HCAPLUS
•	1992	•	2821	J Chem Soc, Perkin T	HCAPLUS
· · · · · · · · · · · · · · ·	•		1377	J Heterocycl Chem	
Isomura, K	1972	1	629		HCAPLUS
Itahara, T	1979		151	. 3	HCAPLUS
Ito, S .	11997		3161	J Chem Soc, Perkin T	HCAPLUS
	11997	119	1486	J Am Chem Soc	HCAPLUS
Jacobi, P	11992.	33	6239	Tetrahedron Lett	HCAPLUS
James, D	11962	27	3346	J Org Chem	HCAPLUS
Jeandon, C	1993		1625	Bull Soc Chim Fr	HCAPLUS
Jefford, C		69	12048	Helv Chim Acta	HCAPLUS
Jefford, C	1996	•	1069	Tetrahedron: Asymmet	HCAPLUS
Johnson, A		i	1950	J Chem Soc C	İ
Jones, G		2	1	Comprehensive Hetero	HCAPLUS
Jones, R	1977	<del>-</del>	1	The Chemistry of Pyr	
•		49	4203	·	HCAPLUS
Kaiser, H		148	3214	-	HCAPLUS
Kakushima, M	•	140		J Chem Soc, Perkin T	•
Kamigata, N	11994	104	11339		HCAPLUS
Kamigata, N	=	124	12049	• 4	
± ,	-	4 4	1374		HCAPLUS
Kashima, C	•		162	·	HCAPLUS
Kato, T	1971	119	1292	1	HCAPLUS
Katritzky, A	1997.		67	· _	HCAPLUS
Katritzky, A	-	34	11379	-	HCAPLUS
Katritzky, A	1993	58	1987	J Org Chem	HCAPLUS
Katritzky, A	•	159	45 <u>.</u> 51	J Org Chem	HCAPLUS
Katritzky, A	1996	61	1624	J Org Chem	HCAPLUS
Katritzky, A	11997	62	14148	J Org Chem	HCAPLUS
Katritzky, A	1988	120	585	Org Prep Proced Int	HCAPLUS
Katritzky, A	1991	1	1863	Synthesis	1
Katritzky, A	1994	1	193	Synthesis	HCAPLUS
Katritzky, A	11995	i	11315	Synthesis	HCAPLUS
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Katritzky, A	1986	42	1623		HCAPLUS
Katritzky, A	11992		•		HCAPLUS
Katritzky, A			•	·	HCAPLUS
Katritzky, A		136		·	HCAPLUS
	1977			•	HCAPLUS
	1980	136			HCAPLUS
Keating, T	11996	118	12574		HCAPLUS
Kelly, T	11993	•		•	HCAPLUS
Ketcha, D	1997	19		Progress in Heterocy	
Ketcha, D	1998	10	109	Progress in Heterocy	HCAPLUS
Khetan, S		124	1567	Tetrahedron	1
Khetan, S		125 -		•	HCAPLUS
Kiely, J			-	· -	HCAPLUS
Kim, I	11998	139		Tetrahedron Lett	HCAPLUS
Kleinspehn, G	1955	177	•		HCAPLUS
Kochhar, K	1984	49			HCAPLUS
Komatsu, M	1993	58			HCAPLUS
Konakahara, T	11993	35	1171	Heterocycles	HCAPLUS
Konieczny, M	11992	133	16939	Tetrahedron Lett	
Korakas, D	1994	1	164	Synthesis	HCAPLUS
Korostova, S	1985	21		J Org Chem USSR (Eng	
Korostova, S	1985	21	•		HCAPLUS
Korostova, S	1984	20	1790	Zh Org Khim, Org Che	
Kost, A	11956	126	1565		HCAPLUS
Kuser, P	1971	54	1969	Helv Chim Acta	HCAPLUS
Kusumoto, T	1986	27	4197	Tetrahedron Lett	HCAPLUS
La Porta, P	1994	1 .	287	Synthesis	HCAPLUS
Lamon, R	1969	6	261	J Heterocycl Chem	HCAPLUS
Lash, T	1991	128	1671	J Heterocycl Chem	HCAPLUS
Lash, T	1993	130	525	J Heterocycl Chem	HCAPLUS
Lash, T	1994	1	170	Synthesis	HCAPLUS
Lash, T		35	2493	Tetrahedron Lett	HCAPLUS
Lash, T		138	2031	•	HCAPLUS
Laurent, A	1982	123	1655	Tetrahedron Lett	HCAPLUS
Lavilla, R	1997	153		Tetrahedron	HCAPLUS
Lee, C	1978	143	3727	J Org Chem	HCAPLUS
Liddell, P	1993	49	1343	Tetrahedron	HCAPLUS
Likhitwitayawuid, K	1987	43	13689	Tetrahedron	HCAPLUS
Madsen, J	1968	24	13369	Tetrahedron	HCAPLUS
Magedov, I	1995	136	1.4619	Tetrahedron Lett	HCAPLUS
Makhsumov, A	1970	16	120	Chem Heterocycl Comp	
Makhsumov, A	1970	16	1393	J Org Chem USSR (Eng	
Makhsumov, A	1970	16	125	Khim Geterotsikl Soe	
Makhsumov, A	11970	16	401	Zh Org Khim	HCAPLUS
Mamedov, E	11983	19	11243	Chem Heterocycl Comp	
Mamedov, E	11983	19	11561	Khim Geterotsikl Soe	!
Mandell, L	11965	2	479	J Heterocycl Chem	
Martina, S	11991	41	403	Synth Met	HCAPLUS
Martina, S	1991		613	Synthesis	HCAPLUS
Maryanoff, B	1977	14	1177	J Heterocycl Chem	HCAPLUS
Maryanoff, B	1979	4 4	14410	J Org Chem	HCAPLUS
Masquelin, T	11995	1	1276	Synthesis	HCAPLUS
Mataka, S	11982	161	157	Synthesis	HCAPLUS
McLeod, M	1996	61	11180	J Org Chem	HCAPLUS
Mendez, J	11996	37	14099	Tetrahedron Lett	HCAPLUS
Merah, B	11980	I .	552 ·	Bull Soc Chim Fr	HCAPLUS
Messinger, P	11986	1	213	Synthesis  Synthesis	HCAPLUS
Meunier, A	11988	1	381	<del>-</del>	HCAPLUS
Meyer, H.	11981	100	11534	Liebigs Ann Chem	HCAPLUS
Middleton, W	1958	180	12822	J Am Chem Soc	HCAPLUS

Mikhlina F	1984	120	1149	Khim Geterotsikl Soe	ı .
Mikhlina, E	11981		5319		HCAPLUS
Minato, A				•	•
Minguez, J	•		4263		HCAPLUS
Mondelli, R	•		17		HCAPLUS
Montforts, F	•		2301		HCAPLUS
Monti, D			1587	·	HCAPLUS
Morgan, K			1245	• = = = = = = = = = = = = = = = = = = =	HCAPLUS
Moskal, J	•		4131		HCAPLUS
Motekaitis, R			12504		HCAPLUS
·	-		1168		HCAPLUS
Muchowski, J	•		1863	_	HCAPLUS
Mukaiyama, T	-		203	. 3	HCAPLUS
Murahashi, S	1974		931	J Chem Soc, Chem Com	HCAPLUS
Muratake, H	1996	4 4	67	Chem Pharm Bull	HCAPLUS
Nagafuji, P	1996	61	14999	J Org Chem	HCAPLUS
Nakajima, S	1995	136	18457	Tetrahedron Lett .	HCAPLUS
Nayyar, N	1997	62	982	J Org Chem	HCAPLUS
Negro, A	1988	120	414	Org Prep Proced Int	HCAPLUS
Neier, R	1996	12	35	Adv Nitrogen Heteroc	
Nenajdenko, V	1997	-	1349	_	HCAPLUS
Nygaard, L	•	•	491		HCAPLUS
Oda, K	-		463	•	HCAPLUS
Okamoto, S	•		6984		HCAPLUS
Ono, N			4470	·	HCAPLUS
Ono, N			3386		HCAPLUS
•	-	•	1707	<del>-</del>	HCAPLUS
Ono, N		44	12020	<del>-</del>	HCAPLUS
Ozaki, S			1813		HCAPLUS
Padwa, A	•	-	-	• • •	HCAPLUS
Padwa, A	-	197	4682	-	HCAPLUS
Padwa, A	•	14	1118	J Heterocycl Chem	•
Padwa, A	•	143	1381	J Org Chem	HCAPLUS
Padwa, A	-	150	14006	J Org Chem	HCAPLUS
Padwa, A	11979		1107	Tetrahedron Lett	HCAPLUS
Padwa, A		129	3041	Tetrahedron Lett	HCAPLUS
Paine, J		150	15598	J Org Chem	HCAPLUS
Paine, J	•	52	13986	J Org Chem	HCAPLUS
Papadopoulos, E	1968	1	1721	Tetrahedron Lett	
Parvi, N	1997	62	2649	J Org Chem	· ·
Patterson, J	1975	197	360	J Am Chem Soc	HCAPLUS
Patterson, J	1968	33	2057	J Org Chem	HCAPLUS
Pedersen, C	1973	27	1271	Acta Chem Scand	HCAPLUS
Pelkey, E	1997	1	1873	Chem Commun (Cambrid	HCAPLUS
Petruso, S	1990	127	1209	J Heterocycl Chem	HCAPLUS
Petruso, S	11990	127	1277	J Heterocycl Chem	HCAPLUS
Petruso, S	11994	31	941	J Heterocycl Chem	
Pfeiffer, G	11976	İ	383	Justus Liebigs Ann C	HCAPLUS
Pfoertner, K		163	658	Helv Chim Acta	HCAPLUS
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Pinna, G	1993	i	1210	J Chem Res	HCAPLUS
Pizzorno, M		142	1909	J Org Chem	HCAPLUS
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Plieninger, H	11970	i	587	Synthesis	HCAPLUS
Prasad, A		53	16711	Tetrahedron	HCAPLUS
Prasad, J		129	14253	Tetrahedron Lett	HCAPLUS
Qui, Z	11994	135	4319	Tetrahedron Lett	1
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Rio, G	11974	1	12824	•	HCAPLUS
Rokach, J		22	14901	•	HCAPLUS
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Roomi, M		180	1702	Angew Chem	I IICAI BOD
Rosenmund, P		17	1733	Angew Chem Int Ed En	I UCADITIC
Rosenmund, P	•	154	14736		HCAPLUS
Roskamp, E		1304	173	- · · · · · · · · · · · · · · · · · · ·	HCAPLUS
Roth, H		311	1844	J Prakt Chem	I IICAI BOS
Ruhlmann, K	11971	1	1236	Synthesis	1
Ruhlmann, K		127	12857	· •	  HCAPLUS
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Sammes, M	11902	160	14947	<del>-</del>	HCAPLUS
Santiago, B		15	119	Chem Res Tox	
Sayre, L Schloemer, G		159	15230	•	  HCAPLUS
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Schmitz, E	•	196	11470	Chem Ber	HCAPLUS
Schulte, K		198	198	Chem Ber	HCAPLUS
Schulte, K Schultz, G	11903	189	1256	Angew Chem	1
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Schultz, G Schumacher, D	11981	146	15060	J Org Chem	HCAPLUS
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Sheradsky, T	11970		25	•	HCAPLUS
Siedel, W		554	1162	Justus Liebigs Ann C	
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Smith, L	11969	34	633	J Org Chem	HCAPLUS
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Stapfer, C	11970	7	651	J Heterocycl Chem	HCAPLUS
Stetter, H	1979	16	839	J Heterocycl Chem	HCAPLUS
Stork, G	1970	1	1445	J Chem Soc D	HCAPLUS
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Tanaseichuk, B		5	1144	Zh Org Khim	HCAPLUS
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Tani, M		44	148	Chem Pharm Bull	HCAPLUS
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Tramontini, M	11973		703	Synthesis	HCAPLUS
Tramontini, M	11990	146	1791	Tetrahedron	HCAPLUS
Tratmer, R	11974	111	189	J Heterocycl Chem	i
Treibs, A	11957	90	179	Chem Ber	HCAPLUS
Treibs, A		1577	105	Justus Liebigs Ann C	
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Zbiral, E
                       HCAPLUS COPYRIGHT 2006 ACS on STN
L32 ANSWER 13 OF 67
                          2001:194786 HCAPLUS Full-text
ACCESSION NUMBER:
                          134:237385
DOCUMENT NUMBER:
```

TITLE:

Preparation of pyrrolidines and their use as

herbicides

INVENTOR(S):

Kato, Masahiko; Yamada, Yasuo; Sato,

Atsushi; Takahashi, Akihiro Nippon Soda Co., Ltd., Japan

PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 24 pp.

SOURCE:

CODEN: JKXXAF

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 2001072661 PRIORITY APPLN. INFO.:	A	20010321	JP 1999-250404 JP 1999-250404	19990903 19990903
OTHER SOURCE(S): GI	CASRE	ACT 134:2373	85; MARPAT 134:237385	

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Pyrrolidines I [A = H, C1-6 (halo)alkyl, C2-6 alkenyl, (un)substituted PhCH2, C1-6 alkylsulfonyl, etc.; R1 = H, C1-6 (halo)alkyl, C2-6 alkenyl, C3-6 cycloalkyl, (un)substituted PhCH2, etc.; R2, R3 = H, C1-6 alkyl; R2R3 may form ring; X = halo, NO2, C1-6 (halo)alkyl, C1-6 alkoxy; n = 0-5] are prepared by cyclocondensation of benzene derivs. II (X, n, R1-R3 = same as above; R4 = C1-6 alkyl), followed by optional modification of the resulting products I (A = H; R1-R3, X, n = same as above). Thus, Et 2-methyl-2-[N-(2,4,6-trimethylphenylacetyl)methoxyamino] propionate was refluxed with Me3COK in THF for 10 min to give 84% I (A = H, R1 = R2 = R3 = Me, Xn = 2,4,6-Me3), which at 2000 g/ha showed 100% herbicidal activity on Digitaria adscendens and Setaria faberi.

L32 ANSWER 14 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:504598 HCAPLUS Full-text

DOCUMENT NUMBER:

CORPORATE SOURCE:

135:242097

TITLE:

Vinyl sulfones in solid-phase synthesis: preparation

of 4,5,6,7-tetrahydroisoindole derivatives

AUTHOR(S):

Cheng, Wei-Chieh; Olmstead, Marilyn M.; Kurth, Mark J.

Department of Chemistry, University of California,

Davis, CA, 95616-5295, USA

SOURCE:

Journal of Organic Chemistry (2001), 66(16), 5528-5533

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal English

LANGUAGE: OTHER SOURCE(S):

CASREACT 135:242097

The preparation of functionalized 4,5,6,7-tetrahydroisoindole via a traceless solid-phase sulfone linker strategy is described. Thermolytic extrusion of SO2 from polymer-bound 3-(phenylsulfonyl)-3-sulfolene generated polymer-bound 2-(phenylsulfonyl)-1,3-butadiene in situ, which underwent Diels-Alder cycloaddn. with various dienophiles to furnish vinyl sulfone resins. To complete a traceless linker cleavage strategy, (p-tolysulfonyl)methyl

isocyanide or Et isocyanoacetate was employed to react with the vinyl sulfone moiety to liberate functionalized 4,5,6,7-tetrahydroisoindole products from the resin. Using this chemical, nine tetrahydroisoindole derivs. were prepared in 32-41% overall yields from polystyrene/divinylbenzene sulfinate. 190449-12-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and solid-phase Diels-Alder reactions of resin-bound (phenylsulfonyl)sulfolene derivs. to isoindole derivs.)

RN 190449-12-8 HCAPLUS

IT

CN

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE Referenced Author (RAU)	Year  (RPY)	(RVL)	(RPG)	Referenced Work   (RWK)	Referenced   File
Abel, Y		81	1978	Helv Chim Acta	HCAPLUS
Andell, O	1985	126	14555	•	HCAPLUS
Ando, K	1994	•	741	•	HCAPLUS
Back, T	•	•	3249	, 5	HCAPLUS
Backvall, J		•	2291	Chem Rev	1
Backvall, J	-	•	6396	J Am Chem Soc	
Backvall, J		•	5221	J Org Chem	
Backvall, J			1445	Tetrahedron Lett	I
Bailey, W	•		1932	•	HCAPLUS
Cheng, W		•	8557	1	HCAPLUS
Chou, S		•	149	•	HCAPLUS
Chou, S		-	1270		HCAPLUS
Chou, T		52	3394		HCAPLUS
Chou, T	•	53	3020		HCAPLUS
Crowley, J		19	135	•	HCAPLUS
Dolle, R		1	235		HCAPLUS
Ellman, J	•		132	Acc Chem Res	HCAPLUS
Farrall, M			3877	J Org Chem	l
Franzen, R		•	195	• • • • • • •	HCAPLUS
Fyles, T	•	•	1935		HCAPLUS
Fyles, T	•		1031	•	HCAPLUS
Haake, G		•	19703	•	HCAPLUS
Halm, C	•	138	7709	•	HCAPLUS
Hoffmann, H		46	5591	•	HCAPLUS
Hopkins, P	•	•	1209	J Org Chem	1
Hoppe, D		13	1789	Angew Chem, Int Ed E	
Inomata, K	•	51	3341	•	HCAPLUS
Kantorowski, E	-	162	16797		HCAPLUS
Lorsbach, B		199	1549	•	HCAPLUS
Padwa, A	,	54	14232	. 3	HCAPLUS
Padwa, A	•	56	12713	•	HCAPLUS
Padwa, A	11992	57	13540	J Org Chem	HCAPLUS

Patchornik, A	1970  92	17587	J Am Chem Soc	HCAPLUS
Portevin, B	12000   43	4582	J Med Chem	HCAPLUS
Saddler, J	1981  103	2110	J Am Chem Soc	HCAPLUS
Sammelson, R	2001  101	137	Chem Rev	HCAPLUS
Schmidt, W	1997	1903	Synlett '	HCAPLUS
van Leusen, A	1972  52	15337	Tetrahedron Lett	1
Vicente, M	1997  38	13639	Tetrahedron Lett	HCAPLUS
Wang, J	1999  1	1524	J Comb Chem	HCAPLUS
Yan, B	1998  63	155	J Org Chem	HCAPLUS

L32 ANSWER 15 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:126741 HCAPLUS Full-text

DOCUMENT NUMBER:

134:340407

TITLE:

Cycloaddition reaction of [60] fullerene with 3,4-fused

pyrrolo-3-sulfolenes

AUTHOR(S):

Ishida, Hiroshi; Itoh, Kenji; Ito, Satoshi; Ono,

Noboru; Ohno, Masatomi

CORPORATE SOURCE:

Department of Molecular Design and Engineering, Graduate School of Engineering, Nagoya University,

Nagoya, 464-8603, Japan

SOURCE:

Synlett (2001), (2), 296-298 CODEN: SYNLES; ISSN: 0936-5214

Georg Thieme Verlag

PUBLISHER: DOCUMENT TYPE:

Journal English

LANGUAGE:

CASREACT 134:340407

OTHER SOURCE(S): Under thermal extrusion of SO2, Et 3,5-dihydro-4,4-dioxo-1H-thieno[3,4c]pyrrole-2-carboxylates cycloadded to C60 to give [60]fullerocyclohexanefused pyrrole-2-carboxylates. While the reaction with N-substituted pyrroles afforded relatively soluble cycloadducts, the isolated solid product from the N-unsubstituted pyrrole was not readily soluble Nevertheless, its structure was confirmed by Boc-protection of the primarily formed cycloadduct. The Bocprotected product could not be obtained directly, yielding the N-unprotected product instead.

190449-12-8 337515-79-4 337515-80-7 IT

337515-81-8

RL: RCT (Reactant); RACT (Reactant or reagent) (cycloaddn. reaction of [60] fullerene with 3,5-dihydro-4,4-dioxo-1Hthieno[3,4-c]pyrrole-2-carboxylates)

RN 190449-12-8 HCAPLUS

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, CN 2,2-dioxide (9CI) (CA INDEX NAME)

337515-79-4 HCAPLUS RN-

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-5-methyl-, ethyl CN ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 337515-80-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-5-(phenylmethyl)-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 337515-81-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4,5(3H)-dicarboxylic acid, 5-(1,1-dimethylethyl) 4-ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE

Referenced Author (RAU)	Year	·	Referenced
Ando, K Guhr, K Hirsch, A McEsen, C Ohno, M Ohno, M Sliwa, W Torres-Garcia, G Tso, H Tsuda, M	1995   51	J Org Chem  J Am Chem Soc  Synthesis  J Am Chem Soc  Synthesis  Tetrahedron  Fullerene Sci Tech  J Org Chem  Tetrahedron Lett  J Chem Soc, Chem Co	HCAPLUS   HCAPLUS   HCAPLUS     HCAPLUS   HCAPLUS   HCAPLUS   HCAPLUS   HCAPLUS
Vicente, M	1997  38  3639	Tetrahedron Lett	HCAPLUS

L32 ANSWER 16 OF 67

HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:607698 HCAPLUS Full-text

DOCUMENT NUMBER:

133:328797

TITLE: Loading a Porphyrin with Fullerene Units AUTHOR(S): Rieder, Alexander; Kraeutler, Bernhard

CORPORATE SOURCE: Institute of Organic Chemistry, University of

Innsbruck, Innsbruck, A-6020, Austria

SOURCE: Journal of the American Chemical Society (2000),

122(37), 9050-9051

CODEN: JACSAT; ISSN: 0002-7863

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

AB One to four C60 fullerene units can be added sequentially to {tetrakis(3,5-ditert-butylphenyl)tetrasulfolenoporphyrinato}zinc by heating in 1,2-

dichlorobenzene. The complexes were characterized based upon FAB mass spectra as well as UV-visible and 1H NMR spectra.

IT 267237-69-4

RL: RCT (Reactant); RACT (Reactant or reagent)
(reactant for stepwise preparation of zinc fullerenoporphyrinate complexes with 1-4 fullerene moieties)

RN 267237-69-4 HCAPLUS

CN Zinc, [[5,11,17,23-tetrakis[3,5-bis(1,1-dimethylethyl)phenyl]7,9,13,15,19,21-hexahydro-1H,3H,25H,27H-tetrathieno[3,4-b:3',4'-g:3'',4''1:3''',4'''-q]porphine-κN25,κN26,κN27,κN28]

2,2,8,8,14,14,20,20-octaoxidato(2-)]-, (SP-4-1)- (9CI) (CA INDEX NAME)

RETABLE Referenced Author (RAU)	(RPY)   (RVL)   (RPG)	Referenced Work
Battersby, A	1994  264  1551	Science   HCAPLUS
Bourgeois, J	1998  81  1835	Helv Chim Acta   HCAPLUS
Chen, C	1996  5  91	Comprehensive Supram HCAPLUS
Crossley, M	1991    1569	· ·
Diederich, F	1999  28  263	Chem Soc Rev   HCAPLUS
Dietel, E	1998    1357	J Chem Soc, Perkin T   HCAPLUS
Dolphin, D	1978	The Porphyrins   .
Drovetskaya, T	1995  36  7971	Tetrahedron Lett   HCAPLUS
Durr, K	1997  130  1375	Chem Ber/Recl   HCAPLUS
Gunter, M	1999    803	J Chem Soc, Chem Com   HCAPLUS
Gust, D	1993  26  198	Acc Chem Res   HCAPLUS
Huber, R	1989  28  848	Angew Chem, Int Ed E

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Imahori, H
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Jentzen, W
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Krautler, B
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Krautler, B
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Krautler, B
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Lehn, J
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Medford, C
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Montforts, F
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Ranasinghe, M
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Rubin, Y
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Scheer, H
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Schuster, D
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Segura, J
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Sessler, J
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Shelnutt, J
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Smith, K
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Waditschatka, R
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Wasielewsky, M
                        |1992 |92
                                     1435
                                            |Chem Rev
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HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 17 OF 67 ACCESSION NUMBER: 2000:748490 HCAPLUS Full-text

DOCUMENT NUMBER:

134:65448

TITLE:

Synthesis of sulfolenobilins and their cyclization

directed to chlorinatozinc-fullerene dyads

AUTHOR(S):

Kutzki, Olaf; Walter, Andreas; Montforts, Franz-Peter

CORPORATE SOURCE: Institut fur Organische Chemie des FB 2 der Universitat Bremen, Bremen, D-28359, Germany SOURCE:

Helvetica Chimica Acta (2000), 83(9), 2231-2245

CODEN: HCACAV; ISSN: 0018-019X Verlag Helvetica Chimica Acta

PUBLISHER: DOCUMENT TYPE: Journal LANGUAGE: German

CASREACT 134:65448 OTHER SOURCE(S):

A novel chlorinatozinc-fullerene dyad was synthesized to model the photosynthetic reaction center. The synthetic key step for the formation of the dyad is an unusual 1-pot reaction of the (sulfolenobilinato)-Ni with concomitant generation of the chlorin macrocycle and linkage to the [5,6]fullerene-C60-Ih. This 1-pot reaction is a complex cascade of single reaction steps with a total yield of 32% and an average yield of 83% for the individual steps. The chlorinatozinc-fullerene dyad is so far one of 3 examples that contain chlorin moieties, the chromophores in naturally occurring photosynthetic systems.

#### 262611-68-7P 313252-68-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reactant for synthesis of sulfolenobilins and their cyclization to chlorinatozinc-fullerene dyads to model the photosynthetic reaction center)

RN 262611-68-7 HCAPLUS

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6-formyl-3,5-dihydro-, CN phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 313252-68-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxaldehyde, 6-bromo-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 190449-12-8

RL: RCT (Reactant); RACT (Reactant or reagent) (reactant for synthesis of chlorinatozinc-fullerene dyads to model the photosynthetic reaction center)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

IT **218628-86-5** 

RL: RCT (Reactant); RACT (Reactant or reagent). (reactant for synthesis of sulfolenobilins and their cyclization to chlorinatozinc-fullerene dyads to model the photosynthetic reaction center)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 190449-12-8DP, reaction products with fullerene-60 313252-73-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (synthesis of sulfolenobilins and their cyclization to chlorinatozinc-fullerene dyads to model the photosynthetic reaction center)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 313252-73-2 HCAPLUS

CN Zinc, [(1,3,7,8-tetrahydro-7,7,12,13,17,18-hexamethyl-22H,24H-thieno[3,4-b]porphine-kN22,kN23,kN24,kN25)

2,2-dioxidato(2-)]-, (SP-4-2)- (9CI) (CA INDEX NAME)

\*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*
RETABLE

Referenced Author (RAU)	·  (RPY)   (RV	L)   (RPG)		Referenced   File
Abel, Y	1998  81		Helv Chim Acta	HCAPLUS
Anon	2000  1-1	0	The Porphyrin Handbo	<b>)</b>
Borovkov, V	1992  65	1533	Bull Chem Soc	HCAPLUS
	1998  81	1835	Helv Chim Acta	HCAPLUS
Cheng, P	1999  89	1	J Chem Soc, Chem Com	ı
Deisenhofer, J	1989  101	872	Angew Chem	HCAPLUS
Deisenhofer, J	1989  28	1829	Angew Chem, Int Ed	1
Deisenhofer, J	1993	1	The Photosynthetic F	<b>?</b>
Diederich, F	1999  32	1537	Acc Chem Res	HCAPLUS <sub>.</sub>
Dietel, E	1998	1981	Chem Commun	HCAPLUS
Drovetskaya, T	1995  36	7971	Tetrahedron Lett	HCAPLUS
Eschenmoser, A	1977  196	1410	Science	HCAPLUS
Gunter, M	1999	1803	Chem Commun	HCAPLUS
Gust, D	1993  26	1198	Acc Chem Res	HCAPLUS
Gust, D	1997  23	621	Res Chem Intermed	HCAPLUS
Gust, D	1991  159	102	Topics in Current Ch	1
Haake, G	1994  35	19703	Tetrahedron Lett	HCAPLUS
Helaja, J	1999	12403	J Chem Soc	1

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Hirsch, A

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Huber, R
                        |1989 |28
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Huber, R
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Imahori, H
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Krautler, B
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Kuciauskas, D
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                                    1929
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Kurreck, H
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Kurreck, H
                                           || J Chem Soc, Chem Com | HCAPLUS
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Marcus, R
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Montforts, F
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Montforts, F
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Montforts, F
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Montforts, F
                        |2000 |112
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Montforts, F
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Montforts, F
Montforts, F
                        |1979 |18
                                     1675
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                        |1981 |20
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Montforts, F
                        |1982 |21
                                    1214
                                            |Angew Chem, Int Ed
Montforts, F
                        |1994 |94
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Montforts, F
                        |1987 |70
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                                            | Helv Chim Acta
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                                                                  IHCAPLUS
Montforts, F
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Tauber, A
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Tkachenko, N
Tome, A
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Vicente, M
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Wasielewski, M
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                        |1999 |
                                     12469
                                           |Chem Commun
Zheng, G
                       HCAPLUS COPYRIGHT 2006 ACS on STN
L32 ANSWER 18 OF 67
                          2000:791082 HCAPLUS Full-text
ACCESSION NUMBER:
                          134:100851
DOCUMENT NUMBER:
                          Diels-alder reaction of tribenzo[b,g,l]thiopheno[3,4-
TITLE:
                          q]porphyrazine as a new path for porphyrazine core
                          modification
                          Nemykin, Victor N.; Polshina, Ann E.; Kobayashi, Nagao
AUTHOR(S):
                          Department of Chemistry, Graduate School of Science,
CORPORATE SOURCE:
                          Tohoku University, Sendai, 980-8578, Japan
                          Chemistry Letters (2000), (11), 1236-1237
SOURCE:
                          CODEN: CMLTAG; ISSN: 0366-7022
                          Chemical Society of Japan
PUBLISHER:
DOCUMENT TYPE:
                          Journal
                          English
LANGUAGE:
                          CASREACT 134:100851
OTHER SOURCE(S):
      Diels-Alder reaction between tribenzo[b,g,l]thiopheno[3,4-q]porphyrazine and
      di-Me acetylenedicarboxylate at ca. 200 °C produced a substituted
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 $\label{tribenzo} tribenzo[b,g,l] porphyrazine \ and \ 3:1 \ unsym. \ substituted \ phthalocyanine, \ which were characterized by UV-VIS, MCD, NMR, and mass spectroscopies.$ 

IT 319925-05-8P

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation, electronic absorption and magnetic CD of core modified porphyrazines via Diels-Alder reaction of tribenzothiophenoporphyrazine with di-Me acetylenedicarboxylate)

RN 319925-05-8 HCAPLUS

CN 1,4-Epithio-29H,31H-tetrabenzo[b,g,l,q]porphyrazine-2,3-dicarboxylic acid, C,C,C-tris(1,1-dimethylethyl)-1,4-dihydro-, dimethyl ester (9CI) (CA INDEX NAME)

PAGE 2-A

3 (D1-Bu-t)

RETABLE Referenced Author (RAU)	Year   VOL  (RPY) (RVL)	(RPG)	Referenced Work   (RWK)	File
Anon Anon Chen, M Cook, M Cook, M Hanack, M Hauschel, B Hedayatulakh, M Kobayashi, N Kobayashi, N Stihler, P	1989- 1-4  1971   5  1989    1997   7  2000   56  2000    1999    1983   296  1990   112  1999   121  1997   130	    1071	Phthalocyanines: Pro  The Chemistry of Syr  J Chem Soc Perkin Tr  J Mater Chem  Tetrahedron  Eur J Org Chem  Eur J Inorg Chem  CR Acad Sci Ser 2  J Am Chem Soc  J Am Chem Soc  Chem Ber	HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS HCAPLUS
Stillman, M	1997  130	1001	Phthalocyanines: Pro	•

L32 ANSWER 19 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2000:853465 HCAPLUS Full-text DOCUMENT NUMBER: 134:175573

Antifungal amides from Piper hispidum and Piper TITLE:

tuberculatum

Navickiene, H. M. D.; Alecio, A. C.; Kato, M. AUTHOR(S):

J.; Bolzani, V. S.; Young, M. C. M.; Cavalheiro,

A. J.; Furlan, M.

CORPORATE SOURCE: Instituto de Quimica, Universidade Estadual Paulista,

Araraquara-SP, 14800-900, Brazil

Phytochemistry (2000), 55(6), 621-626 SOURCE:

CODEN: PYTCAS; ISSN: 0031-9422

Elsevier Science Ltd. PUBLISHER:

DOCUMENT TYPE:

Journal English

LANGUAGE:

GI

Piper hispidum and Piper tuberculatum accumulate amides bearing iso-Bu, AΒ pyrrolidine, dihydropyridone and piperidine moieties. The isolation and characterization of several representatives, including two hitherto unreported amides (I and II), were performed by chromatog. techniques and by anal. of spectroscopic data. The antifungal activity of each amide was determined by direct bioautog. against Cladosporium sphaerospermum.

		L	

Referenced Author (RAU)	(RPY)   (RVL)   (RPG)	
Alecio, A Araujo-Junior, J Bernard, C Duh, C Filho, R Homans, A Kiuchi, F Maxwell, A Miyako, M Parmar, V Rahalison, L Rosario, S	•	Phytochemistry
Shah, S	1986  25  1997	Phytochemistry  HCAPLUS

L32 ANSWER 20 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

2000:132427 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 133:98607

Simple synthesis of a chlorin-fullerene dyad with a TITLE:

novel ring-closure reaction

Montforts, Franz-Peter; Kutzki, Olaf AUTHOR(S):

CORPORATE SOURCE:

Institut fur Organische Chemie der Universitat FB2

Biologie/Chemie, Bremen, 28334, Germany

SOURCE:

Angewandte Chemie, International Edition (2000),

39(3), 599-601

CODEN: ACIEF5; ISSN: 1433-7851

PUBLISHER:

Wiley-VCH Verlag GmbH

DOCUMENT TYPE:

Journal

English

LANGUAGE:

AB

The synthesis of a dyad that contains fullerene C60 bound to chlorin by two methylene bridges by a one-pot reaction is reported. Fluorescence spectra

show that while the bare zinc chlorin complex exhibits luminescence, the luminescence is quenched by the fullerene derivatized zinc chlorin.

262611-68-7P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation and reactant for preparation of zinc chlorin-fullerene dyad)

262611-68-7 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6-formyl-3,5-dihydro-, CN

phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

190449-12-8 ΙT

RL: RCT (Reactant); RACT (Reactant or reagent)

(reactant for preparation of fullerene Diels-Alder adduct)

190449-12-8 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, CN

2,2-dioxide (9CI) (CA INDEX NAME)

218628-86-5

RL: RCT (Reactant); RACT (Reactant or reagent) (reactant for preparation of zinc chlorin-fullerene dyad)

RN 218628-86-5 HCAPLUS

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl

ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)			(RWK)	File
=======================================				+======================================	
Abel, Y	1998		1978	•	HCAPLUS
Armaroli, N	,	•			HCAPLUS
Cheng, P	1999	1	89	J Chem Soc Chem Comm	
Deisenhofer, J	11993			The Photosynthetic R	
Diederich, F	11999		537		HCAPLUS
Dietel, E	1998	•	1981	J Chem Soc Chem Comm	HCAPLUS
Drovetskaya, T	1995	•	•		HCAPLUS
Gust, D	1997	•	•	•	HCAPLUS
Gust, D	1991	1	103	Topics in Current Ch	HCAPLUS
Haake, G	11994	35	19703	•	HCAPLUS
Helaja, J	1999	1	12403	J Chem Soc Perkin Tr	
Hirsch, A	1994	1		The Chemistry of the	
Imahori, H	1997		537		HCAPLUS
Kuciauskas, D	11996	100	115926		HCAPLUS
Kurreck, H	1995	107	1929	Angew Chem	I
Kurreck, H	1995	34	849	Angew Chem Int Ed En	
Linssen, T	1995	Ι,	103	J Chem Soc Chem Comm	HCAPLUS
Martin, N	1998	98	12527	Chem Rev	HCAPLUS
Montforts, F	1981	93	795	Angew Chem	HCAPLUS
Montforts, F	1982	94	1208	Angew Chem	HCAPLUS
Montforts, F	1981	120	778	Angew Chem Int Ed En	1
Montforts, F	11982	21	214	Angew Chem Int Ed En	1
Montforts, F	1982	1	499	Angew Chem Suppl	1
Montforts, F	1994	94	327	Chem Rev	HCAPLUS
Montforts, F	1987	70	402	Helv Chim Acta	HCAPLUS
Montforts, F	1985	1	1228	1 — — — — — — — — — — — — — — — — — — —	HCAPLUS
Montforts, F	1998	10	11	Progress in Heterocy	HCAPLUS
Tkachenko, N	11999	121	19378	•	HCAPLUS
Tome, A	1998	54	11141	Tetrahedron	HCAPLUS
Wasielewski, M	1992	192	435	Chem Rev	HCAPLUS
•					
L32 ANSWER 21 OF 67				006 ACS on STN	
ACCESSION NUMBER:	2000	:21330	1 HCAP	LUS <u>Full-text</u>	
DOCUMENT NUMBER:		342346			••
TITLE:	A sy	mmetri	cal tet	rasulfolenoporphyrin	as reactive
		ding b			
AUTHOR(S):	Krau	tler,	Von Ber	nhard; Sheehan, Craig	S.; Rieder,
		ander			
CORPORATE SOURCE:				nische Chemie, Univer	sitat Innsbruck,
	Inns	bruck,	A-6020	, Austria	
SOURCE:				Acta (2000), 83(3),	583-591
	CODE	N: HCA	CAV; IS	SSN: 0018-019X	•
PUBLISHER:	Verl	ag Hel	vetica	Chimica Acta	
DOCUMENT TYPE:	Jour	nal			•
LANGUAGE:	Germ	an			
				2.	

The efficient preparation of the sym. tetrasulfolenoporphyrin (I; R = 3,5-ditert-butylphenyl) is reported, which shows good solubility in a variety of solvents. In I, 4 sulfone groups that sym. bridge the 8 methylene groups in  $\beta$ -position at the pyrrole rings provide 4 sulfolene (2,5-dihydrothiophene 1,1-dioxide) units which are prone to cleavage by thermally induced extrusion of SO2. I lends itself to specific and multiple refunctionalization by the rèplacement of the sulfone groups by other functionalities in SO2 extrusion/cycloaddn. sequences. The predicted reactivity and the potential of ZnL (H2L = I) as a reactive porphyrin module is tested in exploratory expts. To this end, thermolysis of ZnL in dichlorobenzene solution (at 140°) and in the presence of [5,6]fullerene-C60-Ih gives the (fullereno-porphyrinato)zinc still having 3 sulfolene units and a diagonal (difullerenoporphyrinato)zinc that still has 2 sulfolene units, in 14 and in 43% yield, resp.

IT 267237-70-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cycloaddn. reaction with fullerene C60)

RN 267237-70-7 HCAPLUS

CN Zinc, (N,N-dimethyl-4-pyridinamine-kN1)[[5,11,17,23-tetrakis[3,5-bis(1,1-dimethylethyl)phenyl]-7,9,13,15,19,21-hexahydro-1H,3H,25H,27H-tetrathieno[3,4-b:3',4'-g:3'',4''-1:3''',4'''-q]porphine-kN25,kN26,kN27,kN28] 2,2,8,8,14,14,20,20-octaoxidato(2-)]-, (SP-5-21)- (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

t-Bu

IT 218628-86-5

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation and reactivity of a tetrasulfolenoporphyrin derivative)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 144425-36-5P 267237-67-2P 267237-68-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reactivity of a tetrasulfolenoporphyrin derivative)

RN 144425-36-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 267237-67-2 HCAPLUS

CN 1H, 3H, 25H, 27H-Tetrathieno[3, 4-b:3', 4'-g:3'', 4''-l:3''', 4'''-q]porphine, 5,11,17,23-tetrakis[3,5-bis(1,1-dimethylethyl)phenyl]-7,9,13,15,19,21-hexahydro-, 2,2,8,8,14,14,20,20-octaoxide (9CI) (CA INDEX NAME)

RN 267237-68-3 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 267237-69-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and reactivity of a tetrasulfolenoporphyrin derivative)

RN 267237-69-4 HCAPLUS

CN Zinc, [[5,11,17,23-tetrakis[3,5-bis(1,1-dimethylethyl)phenyl]7,9,13,15,19,21-hexahydro-1H,3H,25H,27H-tetrathieno[3,4-b:3',4'-g:3'',4''1:3''',4'''-q]porphine-κN25,κN26,κN27,κN28]
2,2,8,8,14,14,20,20-octaoxidato(2-)]-, (SP-4-1)- (9CI) (CA INDEX NAME)

RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)	(RVL)	(RPG)	(RWK)	File
=======================================	+====+	-====-	+=====	+==============	
Abel, Y	1998	81	1978	1	HCAPLUS
Ando, K	1995	51	129	Tetrahedron	HCAPLUS
Battersby, A	1994		1551	Science	HCAPLUS
Bourgeois, J	1998	81	1835		HCAPLUS .
Chen, C	1996	5	91	Comprehensive Supram	HCAPLUS
Crossley, M	1991		1569	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	HCAPLUS
Dietel, E	1998		1357	J Chem Soc, Perkin T	HCAPLUS
Dolphin, D	1978			The Porphyrins	1 .
Drovetskaya, T	1995	36	7971	•	HCAPLUS
Durr, K	1997	130	1375	·	HCAPĻUS
Gunter, M	1999		1803	Chem Commun	HCAPLUS
Gust, D	1993	26	198	Acc Chem Res	HCAPLUS
Huber, R	1989	101	1849	Angew Chem	HCAPLUS
Huber, R	1989	28	848	Angew Chem, Int Ed	1
Imahori, H	1995		1265	Chem Lett	HCAPLUS
Isaacs, L	1993	76	1231	Helv Chim Acta	HCAPLUS
Krautler, B	1987	41	277	Chimia	
Krautler, B	1993	176	11626	Helv Chim Acta	I
Kuciauskas, D	11999	121	18604		HCAPLUS
Lehn, J	1995	l ,	1.	Supramolecular Chemi	1
Lidell, P	11997	119	1400	J Am Chem Soc	1
Lidell, P	11994	160	1537	Photochem Photobiol	1
Lindsey, J	11987	52	1827	J Am Chem Soc	HCAPLUS
Linssen, T	11995	l	103	•	HCAPLUS
Newman, M	11972	137	4468	J Org Chem	HCAPLUS
Nierengarten, J	11998	1110	12037	Angew Chem	
Nierengarten, J	11998		1934	Angew Chem, Int Ed	HCAPLUS
Nierengarten, J	1998	1	1545	Chem Commun	HCAPLUS
Ranasinghe, M	1996	137	14797	Tetrahedron Lett	HCAPLUS
Rubin, Y	1993	115	1344	J Am Chem Soc	HCAPLUS
Scheer, H	1991	I	1	Chlorophylls	· ·
Sessler, J	1996	4	311	Comprehensive Supram	HCAPLUS
Smith, K	11975	1	1	Porphyrins and Metal	1
Vicente, M	11997	138	3639	Tetrahedron Lett	HCAPLUS
Wasielewsky, M	11992	92	435	Chem Rev	
Whitlock, H	11969	191	17485	J Am Chem Soc	HCAPLUS

L32 ANSWER 22 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:88097 HCAPLUS Full-text

DOCUMENT NUMBER: 134:274966

TITLE: Synthesis and properties of novel porphine-fullerene

dyads for the investigation of light induced energy

and electron transfer

AUTHOR(S): Kutzki, Olaf; Wedel, Michael; Montforts, Franz-Peter;

Smirnov, Sergei; Cosnier, Serge; Walter, Andreas

CORPORATE SOURCE: University of Bremen, FB 2 - Chemistry/Biology,

Bremen, D-28334, Germany

SOURCE: Proceedings - Electrochemical Society (2000),

2000-10 (Fullerenes 2000--Volume 8: Electrochemistry

and Photochemistry), 172-181 CODEN: PESODO; ISSN: 0161-6374

PUBLISHER: Electrochemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 134:274966

AB A variety of dyads and more complex mol. arrangements of donor and acceptor subunits were designed to investigate light induced energy and electron transfer to mimic natural photosynthesis. Since the discovery and availability of the fullerenes, especially fullerene C60 was used as a subunit for the construction of mol. dyads which undergo light induced electron transfer from a porphinoid donor to the fullerene acceptor. A possible advantage of fullerenes over quinones, which were used as acceptors by nature and in the majority of artificial photosynthetic systems, is the ability of C60 to accept up to six electrons, and is the lower reorganization energy of C60 compared to quinones according to Marcus theory.

IT 218628-86-5P 262611-68-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate product in preparation of zinc porphyrin-fullerene dyad)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 262611-68-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 6-formyl-3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)			(RPG)	• •	File
	1998			•	HCAPLUS
	1975	•		. 3	HCAPLUS
	-			Angew Chem Int Ed En	
Bingel, C	1993		-	•	HCAPLUS
Cosnier, S	1998			J Porphyrins Phthalo	
Diederich, F	1999			•	HCAPLUS
Dinello, R	1978			• •	
Gust, D	1993		•	•	HCAPLUS
Gust, D	11997			Res Chem Intermed	
Gust, D	11991			Topics in Current Ch	
Haake, G	1994	-		•	HCAPLUS
Helaja, J	1999	•		J Chem Soc Perkin Tr	
Hooz, J	1968	-		•	HCAPLUS
Imahori, H	1997		1537	•	HCAPLUS
Imahori, H	1996	-			HCAPLUS
Imahori, H	1999			· •	HCAPLUS
Kurreck, H	1995			1	 
Kurreck, H	1995	•		Angew Chem Int Ed En	
Marcus, R	1993			•	HCAPLUS
Marcus, R	1993	•		Angew Chem Int Ed En	
Martin, N	1998		•		HCAPLUS
Monforts, F	11982	-		Angew Chem Int Ed En	
Montforts, F	1981		795  208	Angew Chem  Angew Chem	HCAPLUS
Montforts, F	11982		1612	Angew Chem	I IICAI IIOS
Montforts, F	2000  1981		1778	Angew Chem Int Ed En	1
Montforts, F	12000		1599	Angew Chem Int Ed En	
Montforts, F	11982		1499	Angew Chem Suppl	I IIOM BOO
Montforts, F Montforts, F	11987		1402		HCAPLUS
Montforts, F	11985		11228	·	HCAPLUS
Montforts, F	1998		1	Progress in Heterocy	•
Nierengarten, J	11998		2037	Angew Chem	
Nierengarten, J	11998		11934	Angew Chem Int Ed En	HCAPLUS
Sakata, Y	11997			_	HCAPLUS
Schumm, O	1928		11	Hoppe-Seyler's Z Phy	
Tkachenko, N				J Am Chem Soc	
Wagner, A					HCAPLUS
Wasielewski, M				•	HCAPLUS
Wedel, M					HCAPLUS
Zehnder, B	11982			Thesis ETH Zurich	İ
Zheng, G	11999		2469	J Chem Soc Chem Comm	HCAPLUS
				1006 - 66 OFW	
				2006 ACS on STN	
ACCESSION NUMBER:				PLUS <u>Full-text</u>	
DOCUMENT NUMBER:		129799			1
TITLE:				nes from porphyrin-fu	
	•			satile porphyrin diene	s for
		oaddit		r	
AUTHOR(S):				J.; Tang, Hesheng	a. Pagland
CORPORATE SOURCE:				stry, University of N	ew England,
COURCE				Australia	9991 (91
SOURCE:	803-		Onunun10	cations (Cambridge) (1	JJJ
	003-	004			

CODEN: CHCOFS; ISSN: 1359-7345

PUBLISHER:

Royal Society of Chemistry

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 131:129799

A porphyrin with a  $\beta$ -fused 3-sulfolene on one of the pyrroles acts as a porphodimethylidene precursor which can be used for a variety of Diels-Alder cycloaddn. reactions.

144425-36-5P 218628-86-5P 234096-95-8P IT

234096-96-9P 234437-54-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of porphodimethylidenes from porphyrin-fused 3-sulfolenes for Diels-Alder cycloaddns.)

144425-36-5 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME) CN

218628-86-5 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl CN ester, 2,2-dioxide (9CI) (CA INDEX NAME)

234096-95-8 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-4-iodo-, 2,2-dioxide (9CI) (CA INDEX CN NAME)

234096-96-9 HCAPLUS RN

1H-Pyrrole-2-carboxylic acid, 5,5'-[(3,5-dihydro-2,2-dioxido-1H-thieno[3,4-CN c]pyrrole-4,6-diyl)bis(methylene)]bis[4-ethyl-3-methyl-, bis(phenylmethyl) ester (9CI) (CA INDEX NAME)

RN 234437-54-8 HCAPLUS

CN 22H,24H-Thieno[3,4-b]porphine, 7,18-diethyl-1,3-dihydro-8,17-dimethyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE					
Referenced Author	Year	VOL	PG	Referenced Work	Referenced
(RAU)	(RPY)	(RVL)	(RPG)	(RWK)	File
=======================================	+====-	+=====	+=====	+======================================	+========
Ando, K	1994		741	Synlett	HCAPLUS
Arnold, D	1994	47	969	Aust J Chem	HCAPLUS
Atkinson, E	1993	34	6147	Tetrahedron Lett	HCAPLUS
Atwood, J	1996	12, 4,		Comprehensive Supram	1
Barton, D	1985	l	1098	J Chem Soc Chem Comm	HCAPLUS
Barton, D	1990	46	17587	Tetrahedron	HCAPLUS
Charlton, J	1987	43	2873	Tetrahedron (Tetrahe	HCAPLUS
Chen, H	1998	54	609	Tetrahedron	
Chou, S	1990	131	1035	Tetrahedron Lett	HCAPLUS
Chou, T	11993	18	65	Rev Heteroatom Chem	HCAPLUS
Crossley, M	11995	l	1921	$ {\tt J}\ {\tt Chem}\ {\tt Soc}\ {\tt Chem}\ {\tt Comm}$	HCAPLUS
Crossley, M	1995	l	2379	$\mid$ J Chem Soc Chem Comm	HCAPLUS
Crossley, M	1996		12675	J Chem Soc Perkin Tr	HCAPLUS
Crossley, M	1997	38	6751	Tetrahedron Lett	HCAPLUS
Faustino, M	11996	37	3569	Tetrahedron Lett	HCAPLUS
Gulyas, P	1997	62	3038	J Org Chem	HCAPLUS
Hopkins, P	•	43	1208	J Org Chem	HCAPLUS
Ito, S	1998	1 .	1661	Chem Commun	HCAPLUS
Ito, S	1997	1	3161	J Chem Soc Perkin Tr	HCAPLUS

Jolliffe, K	1998  37	1916	Angew Chem Int Ed	HCAPLUS
Kai, S	1996  37	5931	Tetrahedron Lett	HCAPLUS
Knapp, S	1998  37	12368	Angew Chem Int Ed	HCAPLUS
Lash, T	1996  2	1197	Chem Eur J	HCAPLUS
Liddell, P	1994  35 -	1995	Tetrahedron Lett	HCAPLUS
Tome, A	1997	11199	Chem Commun	HCAPLUS
Vicente, M	1998	2355	Chem Commun	HCAPLUS
Vicente, M	1997  38	13639	Tetrahedron Lett	HCAPLUS
Warrener, R	1990	1	J Org Chem in press	1
Warrener, R	1998	1590	Synlett	HCAPLUS
Warrener, R	1998	1593	Synlett	HCAPLUS
Zheng, G	1996	1119	Chem Lett	HCAPLUS

L32 ANSWER 24 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN 1998:674757 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 129:349077

TITLE: Ink-jet printing sheet

Kaneko, Akira; Nakamura, Takayuki INVENTOR(S): Mitsubishi Paper Mills, Ltd., Japan PATENT ASSIGNEE(S):

Jpn. Kokai Tokkyo Koho, 17 pp.

CODEN: JKXXAF

Patent DOCUMENT TYPE: LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

SOURCE:

PATENT NO.	KIND	DATE ·	APPLICATION NO.	DATE
JP 10278415	Α	19981020	JP 1997-89447	19970408
PRIORITY APPLN. INFO.:			JP 1997-89447	19970408

In the title printing sheet having an ink-receiving layer, the ink-receiving layer is formed by a gelatin-based graft copolymer. The ink-receiving layer may further contain a hydrophilic vinyl polymer, a fluorine surfactant, polymer fine particles or inorg. particles. The invention printing sheet can be used for high quality color printing with nice gloss.

L32 ANSWER 25 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN 1998:680468 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER:

130:110096 Cruciform porphyrin pentamers TITLE:

Vicente, M. Graca H.; Cancilla, Mark T.; Lebrilla, AUTHOR(S):

Carlito B.; Smith, Kevin M.

Department of Chemistry, University of California, CORPORATE SOURCE: .

Davis, CA, 95616, USA

Chemical Communications (Cambridge) (1998), (21), SOURCE:

2355-2356

CODEN: CHCOFS; ISSN: 1359-7345

DOCUMENT TYPE:

Royal Society of Chemistry Journal

LANGUAGE:

PUBLISHER:

English

OTHER SOURCE(S):

CASREACT 130:110096

GΙ

<sup>\*</sup> STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

AB Cruciform porphyrin pentamers (I) are obtained in good yield by acid-catalyzed tetramerization of ZnII-pyrroloporphyrin (II), followed by oxidation with DDQ; pyrroloporphyrins are in turn obtained from the corresponding pyrrolochlorins by Diels-Alder type reactions of porphyrins involving thermal extrusion of sulfur dioxide from a pyrrole-fused 3-sulfolene.

IT 190449-12-8

RL: RCT (Reactant); RACT (Reactant or reagent)
 (synthesis of cruciform porphyrin pentamers)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE Referenced Author	Year   VOL	l PG	Referenced Work   Referenced
(RAU)	(RPY)   (RVL)		,
=======================================	=+=======	+=====	=+=====================================
Clement, T	1998  37	1150	Inorg Chem   HCAPLUS
Crossley, M	1991	1569	J Chem Soc Chem Comm   HCAPLUS
Crossley, M	1995	1921	J Chem Soc Chem Comm   HCAPLUS
Crossley, M	1995	2379	J Chem Soc Chem Comm   HCAPLUS
Crossley, M	1997  38	6751	Tetrahedron Lett   HCAPLUS
Green, M	1997  3	1439	Eur J Mass Spectrom   HCAPLUS
Gros, C	1997  1	201	J Porphyrins Phthalo HCAPLUS
Jaquinod, L	1996  35	1840	Angew Chem Int Ed En HCAPLUS
Jaquinod, L.	1996	11475	Chem Commun
Jaquinod, L	1998	1261	Chem Commun   HCAPLUS
Khoury, R	1997	1057	Chem Commun   HCAPLUS
Norsten, T	1998	1257	Chem Commun   HCAPLUS
Ono, N	1990  46	17483	Tetrahedron   HCAPLUS
Osuka, A	1997  36	135 `	Angew Chem Int Ed En HCAPLUS
Reek, N	1997  36	361	Angew Chem Int Ed En
Tome, A	1997	1199	Chem Commun   HCAPLUS
Vicente, M	1997  38	3639	Tetrahedron Lett   HCAPLUS
Warrener, R	1998	593	Synlett   HCAPLUS
Zheng, G	1997  38	12409	Tetrahedron Lett     HCAPLUS

L32 ANSWER 26 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1998:756298 HCAPLUS Full-text

130:81364

DOCUMENT NUMBER:

TITLE:

A simple and flexible synthesis of pyrroles from

 $\alpha, \beta$ -unsaturated sulfones

AUTHOR(S): Abel, Yvonne; Haake, Elmar; Haake, Gerold; Schmidt,

Wolfgang; Struve, Daria; Walter, Andreas; Montforts,

Franz-Peter

CORPORATE SOURCE: Institut Organische Chemie, FB 2, Universitaet Bremen,

Bremen, D-28359, Germany

SOURCE: Helvetica Chimica Acta (1998), 81(11), 1978-1996

CODEN: HCACAV; ISSN: 0018-019X

PUBLISHER:

Verlag Helvetica Chimica Acta AG

DOCUMENT TYPE:

Journal

LANGUAGE:

German

OTHER SOURCE(S):

CASREACT 130:81364

AB The addition of alkyl isocyanoacetates and isocyanoacetonitrile to  $\alpha,\beta$ -unsatd. sulfones affords a convenient and broad access to pyrroles with unusual substitution patterns. The  $\alpha,\beta$ -unsatd. sulfones required as starting materials are easily obtained from olefins.

IT **218628-86-5P** 

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of pyrroles from  $\alpha,\beta$ -unsatd. sulfones)

RN 218628-86-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RETABLE Referenced Author	Year	ı vol	l PG	Referenced Work	Referenced
(RAU)		(RVL)		(RWK)	File .
The cluster A	+=====	+====- '	+===== 	+=====================================	
Abdel Hady, A	11997	130	11745	·	IHCAPLUS
Abel, Y	,	•	1741	Synlett	l HCAL BOS
Ando, K Anon	11980	·	1133	Angew Chem Int Ed En	1 1
Anon	•	-	1914	Angew Chem Int Ed En	
Appel, R	11975	•	1863		' .  HCAPLUS
Appel, R		•	1801	Angew Chem Int Ed En	•
		•	1969	Aust J Chem	! 
Arnold, D Barkingia, K		•	17566	J Am Chem Soc	! 
Barkingia, K Barkingia, K	•		18851	IJ Am Chem Soc	I
Barton, D	11985		11098	IJ Chem Soc Chem Comm	IHCAPLUS
Barton, D		•	17587	• •	HCAPLUS
Burns, D	•	•	1379		HCAPLUS
Buschmann, J			1988	Angew Chem	1
			•	Tetrahedron Lett	1
Dolphin, D	•	•	12267	Chem Rev	, 1
Eschenmoser, A	•	•	15	'	,  HCAPLUS
Eschenmoser, A		-	15	Angew Chem Int Ed En	•
Falk, H	11989	_ , 	1	The Chemistry of Lin	
Gossauer, A	11974	i I	i	Die Chemie der Pyrro	
Gust, D	•	145	4867	<u>-</u>	HCAPLUS
Haake, G	11994	•	19703	• • • • • • • • • • • • • • • • • • • •	HCAPLUS
Helmchen, G	•		409		HCAPLUS
Hopkins, P	•	-	1209	J Org Chem	1
Hoppe, D	•	•	1878		HCAPLUS
Hoppe, D		•	789	Angew Chem Int Ed En	i
Houwing, H	•	-	1143	Tetrahedron Lett	
Ito, S	11997	•	3161	J Chem Soc Perkin Tr	HCAPLUS
Jentzen, W	1995	1117	11805	J Am Chem Soc	İ
•	•		•		•

Jones, A	1977	1	The Chemistry of Pyr
Liotta, D	1981  46	12605	J Org Chem     HCAPLUS
Magnus, P	1984  25	1421	Tetrahedron Lett
Montforts, F	1994  94	327	Chem Rev   HCAPLUS
Montforts, F	1998  E9d	577	Houben-Weyl Methods
Montforts, F	1998  10		Progress in Heterocy HCAPLUS
Muller, H	1968  90	12075	J Am Chem Soc
Nicolaou, K	1979  101	13884	J Am Chem Soc   HCAPLUS
Scarborough, R	1977  50	14361	Tetrahedron Lett
Schollkopf, U	1977  89	351	Angew Chem
Schollkopf, U	1977  16	1339	Angew Chem Int Ed En
Scmidt, W	1997	1903	Synlett
Seitz, U	1986	1686	Synthesis   HCAPLUS
Sessler, J	1997		Expanded Contracted
Simpkins, N	1993	1	Sulfones in Organic
Trost, B	1981  22	1287	Tetrahedron Lett   HCAPLUS
van Leusen, A	1972  52	15337	Tetrahedron Lett
van Leusen, A	1975  40	3487	Tetrahedron Lett
Vicente, M	1997  38	13639	Tetrahedron Lett   HCAPLUS
Wilson, S	1980  10	339	Synth Commun   HCAPLUS
Wissman, H	1980  92	129	Angew Chem

L32 ANSWER 27 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1997:349359 HCAPLUS Full-text

DOCUMENT NUMBER:

127:17531

TITLE:

Synthesis and cycloaddition reactions of pyrrole-fused

3-sulfolenes: a new versatile route to

tetrabenzoporphyrins

AUTHOR(S):

Vicente, Maria G. H.; Tome, Augusto C.; Walter,

Andreas; Cavaleiro, Jose A. S.

CORPORATE SOURCE:

Department of Chemistry, University of Aveiro, Aveiro,

3810, Port.

SOURCE:

Tetrahedron Letters (1997), 38(20), 3639-3642

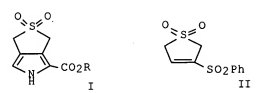
CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S):

CASREACT 127:17531

GΙ



AB Pyrrole-fused 3-sulfolenes I (R = Et, CMe3) were prepared from the corresponding  $\alpha, \beta$ -unsatd. sulfone II. These pyrroles undergo thermal extrusion of sulfur dioxide to produce highly reactive o-quinodimethanes which can be trapped in Diels-Alder reactions. The resulting pyrroles are important starting reagents in porphyrin synthesis.

IT 190449-12-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(preparation of tetrabenzoporphyrins via cycloaddn. of pyrrole-fused sulfolenes)

RN 190449-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, ethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 190449-32-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of tetrabenzoporphyrins via cycloaddn. of pyrrole-fused sulfolenes)

RN 190449-32-2 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-4-carboxylic acid, 3,5-dihydro-, 1,1-dimethylethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

lYear   VOL	i PG	Referenced Work   Referenced
(RPY)   (RVL)	(RPG)	(RWK)   File
1994  37  1992    1994    1995  51  1994  47  1985    1990  46  1987  43  1996  100  1990  31  1993  8  1995    1976  98  1994  35  1978  43  1981	1417  1100  741  129  969  1098  7587  2873  17507  1035  65  2537  7638  9703  1208  1005	J Chem Soc, Chem Com   HCAPLUS   Tetrahedron   HCAPLUS   Tetrahedron   HCAPLUS
1991  23	[237	Organic Prep and ProHCAPLUS
	(RPY)   (RVL) =+====+====  1994   37  1992    1994    1995   51  1994   47  1985    1990   46  1987   43  1996   100  1990   31  1993   8  1995    1976   98  1994   35  1978   43  1981	1994   37

Remy, D	.  1983  24	1451	Tetrahedron Lett	HCAPLUS
Tang, J	1994  59	17793	J Org Chem	HCAPLUS
Tome, A	1996	531	Synlett	HCAPLUS
Tome, A	1996  52	11735	Tetrahedron	HCAPLUS
Vancott, T	1993  97	17417	J Phys Chem	HCAPLUS
Vinogradov, S	1995	103	J Chem Soc Perkin	Tr HCAPLUS
Vogler, A	1978  17	1760	Angew Chem Int Ed	En
Wolford, S	1995  24	52	Fundam Appl Toxico	ol   HCAPLUS

HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 28 OF 67 1998:190795 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER: 128:270553

Synthesis of 4-substituted 3,5-dihydro-1H-thieno[3,4-TITLE:

clpyrrole 2,2-dioxides and their Diels-Alder reactions Suzuki, Takayoshi; Ohyabu, Hiroaki; Takayama, Hiroaki

AUTHOR(S): Fac. Pharmaceutical Sci., Teikyo Univ., Sagamiko,

CORPORATE SOURCE:

Kanagawa, 199-01, Japan

Heterocycles (1997), 46, 199-202 SOURCE: CODEN: HTCYAM; ISSN: 0385-5414

Japan Institute of Heterocyclic Chemistry

DOCUMENT TYPE: Journal English LANGUAGE:

CASREACT 128:270553 OTHER SOURCE(S):

The preparation of 4-substituted 3,5-dihydro-1H-thieno[3,4-c]pyrrole 2,2-AB dioxides and their intermol. Diels-Alder reactions with DMAD are described. In the reaction with DMAD, 4-acetyl-3,5-dihydro-1H-thieno[3,4-c]pyrrole 2,2dioxides acted as the corresponding 3,4-dimethylenepyrroles.

IT 144425-39-8

PUBLISHER:

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation and Diels-Alder reactions of 4-substituted 3,5-dihydro-1Hthieno[3,4-c]pyrrole 2,2-dioxides)

144425-39-8 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-methyl-, 2,2-dioxide (9CI) (CA CN INDEX NAME)

205514-14-3P 205514-15-4P 205514-16-5P IT

205514-17-6P 205514-18-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and Diels-Alder reactions of 4-substituted 3,5-dihydro-1Hthieno[3,4-c]pyrrole 2,2-dioxides)

205514-14-3 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(methylsulfonyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 205514-15-4 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4-bromo-3,5-dihydro-5-(methylsulfonyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 205514-16-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4-acetyl-3,5-dihydro-5-(methylsulfonyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 205514-17-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4-bromo-3,5-dihydro-5-methyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 205514-18-7 HCAPLUS

CN Ethanone, 1-(3,5-dihydro-5-methyl-2,2-dioxido-1H-thieno[3,4-c]pyrrol-4-yl)(9CI) (CA INDEX NAME)

RETABLE

Referenced Author | Year | VOL | PG | Referenced Work | Referenced (RAU) | (RPY) | (RVL) | (RPG) | (RWK) | File

1994  37	1417	Heterocycles
1991	1765	J Chem Soc, Chem Com   HCAPLUS
1992	1100	J Chem Soc, Chem Com   HCAPLUS
1992	870	J Chem Soc, Chem Com   HCAPLUS
1993	2263	J Chem Soc, Perkin T   HCAPLUS
1994	741	Synlett   HCAPLUS
1995  51	129	Tetrahedron   HCAPLUS
1	1	CAChe 3.9 system (CA
1995  117	3405	J Am Chem Soc   HCAPLUS
1993	[2387	J Chem Soc, Perkin T   HCAPLUS
1995  36	1865	Tetrahedron Lett   HCAPLUS
1984  49	2795	J Org Chem ·   HCAPLUS
1991  39	2164	Chem Pharm Bull   HCAPLUS
1993  35		Heterocycles   HCAPLUS
1994  38	1961	Heterocycles   HCAPLUS
1990	1687	J Chem Soc, Chem Com   HCAPLUS
1995	1807	J Chem Soc, Chem Com   HCAPLUS
1996	12699	J Chem Soc, Perkin T   HCAPLUS
1997	251	J Chem Soc, Perkin T  HCAPLUS
1988	1044	J Chem Soc, Chem Com   HCAPLUS
1979	583	Chem Lett
1983	1003	!Chem Lett   HCAPLUS
1983  48	13483	J Org, Chem   HCAPLUS
1986  51	14934	J Org, Chem   HCAPLUS
1981  22	2591	Tetrahedron Lett   HCAPLUS
	1991   1992   1993   1994   1995   1995   1995   1995   1995   1997   1998   1997   1988   1979   1983   1983   1986   51	1991    1765 1992    1100 1992    870 1993    2263 1994    741 1995  51  129       1995  117  3405 1993    2387 1995  36  1865 1984  49  2795 1991  39  2164 1993  35  57 1994  38  961 1990    1687 1995    807 1996    2699 1997    251 1988    1044 1979    583 1983  48  3483 1986  51  4934

L32 ANSWER 29 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1995:856612 HCAPLUS Full-text

DOCUMENT NUMBER:

123:270860

TITLE:

SOURCE:

Treatment of lithographic printing plate with good

printability

INVENTOR(S):
PATENT ASSIGNEE(S):

Kaneko, Akira; Saikawa, Masahiko
Mitsubishi Paper Mills Ltd, Japan

Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE: LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 07199473	Α .	19950804	JP 1994-495	19940107
JP 3301844	B2	20020715		
PRIORITY APPLN. INFO.:			JP 1994-495	19940107

AB The plate comprising a coarsened and anodized Al support coated with a physodevelopment nucleus layer (A) and a photosensitive Ag halide emulsion layer (B) is treated by developing and treating with a solution containing ≥1 kind of alkylene oxides or polyalkylene oxides, where A contains no binders and contacts with the anodized layer or A contains a hydrophilic colloid other than gelatin. The plate shows good ink receptibility and printability.

L32 ANSWER 30 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1996:27220 HCAPLUS Full-text

DOCUMENT NUMBER:

124:201907

TITLE:

Synthesis of furan-, thiophene- and pyrrole-fused

sultines and their application in Diels-Alder

reactions

AUTHOR(S):

Chung, Wen-Sheng; Lin, Wen-Ju; Liu, Wen-Dar; Chen,

Liang-Gyi

CORPORATE SOURCE: Department Applied Chemistry, National Chiao Tung

University, Hsinchu, 30050, Taiwan

SOURCE: Journal of the Chemical Society, Chemical

Communications (1995), (24), 2537-9

CODEN: JCCCAT; ISSN: 0022-4936

PUBLISHER: Royal Society of Chemistry

DOCUMENT TYPE: Journal LANGUAGE: English

AB The synthesis of 1,4-dihydrofurano[3,4-d]-3,2-oxathiin 2-oxide, 5,7-dimethyl-1,4-dihydrothieno[3,4-d]-3,2-oxathiin 2-oxide and 1,4-dihydro-6-

tosylpyrrolo[3,4-d]3,2-oxathiin 2-oxide, precursors for nonclassical heteroarom. o-quinodimethanes, and their application in the Diels-Alder

reactions are reported.

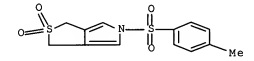
IT 144425-37-6P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 144425-37-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-[(4-methylphenyl)sulfonyl]-,

2,2-dioxide (9CI) (CA INDEX NAME)



L32 ANSWER 31 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1995:542333 HCAPLUS Full-text

DOCUMENT NUMBER: 123:169572

TITLE: Intramolecular Diels-Alder reaction of new building

blocks, N-substituted 3,5-dihydro-1H-thieno[3,4-c]pyrrole S,S-dioxides; a general route to the

tricyclic azanorbornane framework

AUTHOR(S): Suzuki, Takayoshi; Takayama, Hiroaki

CORPORATE SOURCE: Faculty Pharmaceutical Sciences, Teikyo University,

Kanagawa, 199-01, Japan

SOURCE: Journal of the Chemical Society, Chemical

Communications (1995), (8), 807-8 CODEN: JCCCAT; ISSN: 0022-4936

PUBLISHER: Royal Society of Chemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 123:169572

In spite of the absence of activating groups for dienophiles, N-substituted 3,5-dihydro-1H-thieno[3,4-c]pyrrole S,S-dioxides which contain terminal olefin substituents, undergo facile intramol. Diels-Alder reaction and subsequent spontaneous desulfonylation to give the corresponding tricyclic azanorbornane framework in good yields.

IT 167111-12-8P 167111-13-9P 167111-14-0P

167111-15-1P 167111-16-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of tricyclic azanorbornanes by intramol. Diels-Alder reaction of terminal olefin-substituted dihydrothienopyrrole dioxides)

RN 167111-12-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-5(3H)-carboxylic acid, 2-propenyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 167111-13-9 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(1-oxo-4-pentenyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 167111-14-0 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(1-oxo-3-butenyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 167111-15-1 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 5-(3-butenylsulfonyl)-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 167111-16-2 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(2-propenylsulfonyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

L32 ANSWER 32 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:323621 HCAPLUS Full-text

DOCUMENT NUMBER:

122:160510

TITLE:

New building blocks, 3,5-dihydro-1H-thieno-3,4-

c]pyrrole 2,2-dioxides; preparation and their

Diels-Alder reaction with dimethyl

acetylenedicarboxylate

AUTHOR(S):

Ando, Kaori; Kankake, Mutsuo; Suzuki, Takayoshi;

Takayama, Hiroaki

CORPORATE SOURCE:

Fac. Pharm. Sci., Teikyo Univ., Kanagawa, 199-1, Japan

SOURCE: Tetrahedron (1995), 51(1), 129-38

CODEN: TETRAB; ISSN: 0040-4020

PUBLISHER:

Elsevier Journal

DOCUMENT TYPE: LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 122:160510

GI

# \* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

New building blocks, 3,5-dihydro-1H-thieno[3,4-c]pyrrole 2,2-dioxides I (R = CH2Ph, H, 4-MeC6H4SO2, COPh, CO2CH2Ph), have been prepared by the oxidation of their corresponding pyrroline derivs. II with DDQ or Chemical MnO2. The Diels-Alder reaction of I with di-Me acetylenedicarboxylate gave new types of compds.: 7-aza-2,3-dimethylenenorbornenes III, the 1:2 adducts IV, 1a,3a,6,9-tetrahydrobenz[g]indoles V, and dihydroindololsulfolene VI depending on the reaction conditions and the N-substituents. The reaction of I with bis(tert-butylsulfonyl)acetylene was also described.

IT 144425-31-0P 144425-35-4P 144425-37-6P

144425-38-7P 144425-39-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and Diels-Alder reaction of thienopyrrole dioxides)

RN 144425-31-0 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(phenylmethyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

$$0 \longrightarrow N \longrightarrow CH_2 - Ph$$

RN 144425-35-4 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-5(3H)-carboxylic acid, phenylmethyl ester,

#### 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-37-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-[(4-methylphenyl)sulfonyl]-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-38-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 5-benzoyl-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-39-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-methyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 144425-36-5P 161201-58-7P 161201-59-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation and Diels-Alder reaction of thienopyrrole dioxides)

RN 144425-36-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 161201-58-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4-[1,2-bis[(1,1-dimethylethyl)sulfonyl]ethenyl]-3,5-dihydro-5-methyl-, 2,2-dioxide, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 161201-59-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 4,6-bis[1,2-bis[(1,1-dimethylethyl)sulfonyl]ethenyl]-3,5-dihydro-5-methyl-, 2,2-dioxide, (E,E)-

dimethylethyl)sulfonyljethenylj-3,5-dinydro-5-methyl-, 2,2-dioxide, (E,E)(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L32 ANSWER 33 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1995:19187 HCAPLUS Full-text

DOCUMENT NUMBER:

122:92438

TITLE:

Photochromic compound

INVENTOR(S):

Tomoda, Akihiko; Suzuki, Hisao; Kaneko, Akira

; Tsuboi, Hideki

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

U.S., 13 pp. Cont. of U.S. Ser. No. 607,999,

abandoned.
CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

#### PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
				-	
US 5296607	Α	19940322	US 1992-902778		19920623
PRIORITY APPLN. INFO.:			JP 1989-90578	Α	19891108
			JP 1990-89664	Α	19900718
			US 1990-607999	В1	19901031

OTHER SOURCE(S): MARPAT 122:92438 For diagram(s), see printed CA Issue.

The title compds. are described by the general formula I (ring A = II, III, AB IV, or V; R1-4 = H, an alkyl group, or an aryl group; X = O or N-R6; when A =II, R5 = an electron donating group in the form of an indolyl group described by the general formula VI or a group described by the general formula VII; when A = III, IV, or V, R5 = an amino group, a di-alkyl amino group, a monoalkyl amino group, a pyrrolyl group, an indolyl group, VI, or VII; R6 = H, an alkyl group, an aryl group, or an allyl group; R7 = an alkyl group, an aryl group, an aralkyl group, or an allyl group; R8 = an alkoxyl group, an amino group, a di-alkyl amino group, or a mono-alkyl amino group; ring B = a 5 or 6 membered heteroarom. ring or a condensed ring containing a 5 or 6 membered heteroarom. ring; Z = O, S, N-R9; R9 = an alkyl group, an aryl group, an aralkyl group, or an allyl group; k = 1 or 2; for V, k can also be 3 or 4; l =an integer in the range 0-5; m = an integer in the range 1-5; for groups attached to II, m can addnl. be 0; and n = an integer from 0 to 4). The introduction of electron donating groups into the heteroarom. structure is effective in improving the long wave length sensitivity without lowering thermal stability. Application is indicated to optical recording media, copying media, printing media, optical filters, and display materials.

L32 ANSWER 34 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN . 1995:99769 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER:

122:55849

TITLE:

Diels-Alder reaction of 3,5-dihydro-1H-thieno[3,4c]pyrrole 2,2-dioxides with alkene dienophiles; facile

preparation of 4,5,6,7-tetrahydroisoindoles

AUTHOR(S):

Ando, Kaori; Kankake, Mutsuo; Suzuki, Takayoshi;

Takayama, Hiroaki

CORPORATE SOURCE:

Fac. Pharmaceutical Sciences, Teikyo Univ., Sagamiko,

199-01, Japan

SOURCE:

Synlett (1994), (9), 741-2 CODEN: SYNLES; ISSN: 0936-5214

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 122:55849

GΙ

3,5-Dihydro-1H-thieno[3,4-c]pyrrole 2,2-dioxides I (R = CH2Ph, Me, CO2CH2Ph, AB COPh) when heated with alkene dienophiles (di-Me fumarate, di-Me maleate, trans-1,2-bis(phenylsulfonyl)ethylene) sequentially underwent Diels-Alder reaction on the pyrrole moiety, a cheletropic elimination of sulfur dioxide, a

Diels-Alder reaction of the resulting diene, and a retro Diels-Alder reaction to give 4,5,6,7-tetrahydroisoindoles II (R1 = R2 = CO2Me, R3 = H; R1 = R3 = CO2Me, R2 = H; R = CO2CH2Ph, R1 = R2 = SO2Ph, R3 = H) in excellent yields.

IT 144425-31-0 144425-35-4 144425-38-7

144425-39-8

RL: RCT (Reactant); RACT (Reactant or reagent)
 (Diels-Alder of thienopyrrole dioxides with alkene dienophiles in
 preparation of isoindoles)

RN 144425-31-0 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(phenylmethyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-35-4 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-5(3H)-carboxylic acid, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-38-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 5-benzoyl-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

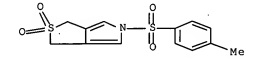
RN 144425-39-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-methyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

RL: RCT (Reactant); RACT (Reactant or reagent) (attempted Diels-Alder with alkene dienophiles in preparation of isoindoles)

144425-37-6 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-[(4-methylphenyl)sulfonyl]-, CN 2,2-dioxide (9CI) (CA INDEX NAME)



L32 ANSWER 35 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1994:548983 HCAPLUS Full-text

DOCUMENT NUMBER:

121:148983

TITLE:

fifty-two-week oral toxicity study of the new

cognition-enhancing agent nefiracetam in dogs

AUTHOR(S):

Hooks, W. N.; Burford, P.; Begg, S.; Gopinath, C.;

Inage, F.; Kato, M.; Takayama, S.

CORPORATE SOURCE:

Huntingdon Res. Cent. Ltd., Huntingdon/Cambridgeshire,

SOURCE:

Arzneimittel-Forschung (1994), 44(2a), 228-38

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE:

Journal

LANGUAGE:

English

A 52-wk toxicity study by oral administration (capsule) was performed in AB beagle dogs with nefracetam (N-(2,6-dimethylphenyl)-2-(2-oxo-I- pyrrolidinyl) acetamide, DM-9384, CAS 77191-36-7), a new cognition-enhancing agent, as a part of a safety evaluation program. Dosage of 0 (control), 10, 30 and 90 mg/kg/d were selected for this study. Treatment-related findings were confined to the 90 mgkg/d level and indicated the kidney and the testis as the main target organs for toxicity. Signs of systemic toxicity, as indicated by the laboratory investigations, were not apparent until the second half of the study and included the principal findings of higher urea nitrogen, and creatinine with higher urinary vols. and corresponding lower sp. gr., osmolarity and protein values. The microscopic pathol. examination showed various changes at the renal papilla, collecting ducts, and medullary and cortical scarring. This examination also revealed decreased spermatogenesis in the testes, with associated decreased nos. or absence of spermatozoa in the epididymides. At the 30 mg/kg/d level, the minor microscopic pathol. changes seen in the kidneys of one male animal were considered to be of equivocal toxicol. importance. There were no treatment-related findings at the low dosage level (10 mg/kg/d) and, therefore, this level was considered as the non-toxic effect level of nefiracetam.

L32 ANSWER 36 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER:

1994:548982 HCAPLUS Full-text

DOCUMENT NUMBER:

121:148982

TITLE:

Fifty-two-week oral toxicity study of the new cognition-enhancing agent nefiracetam in rats

AUTHOR(S):

Hooks, W. N.; Colman, K. A.; Gopinath, C.; Inage, F.;

Kato, M.; Takayama, S.

CORPORATE SOURCE:

Huntingdon Res. Cent. Ltd., Huntingdon/Cambridgeshire,

UK

SOURCE: Arzneimittel-Forschung (1994), 44(2a), 220-8

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE: Journal LANGUAGE: English

A 52-wk toxicity study by oral gavage administration was performed in Sprague-Dawley rats with nefiracetam (N-(2,6-dimethylphenyl)-2-(2-oxo-1pyrrolidiny1) acetamide, DM-9384, CAS 77191-36-7), a new cognition-enhancing agent, as a part of a safety evaluation program. Dosages of 0 (control), 10, 30, 100 and 300 mg/kg/d were selected for this study. Treatment-related findings were confined to the 300 mg/kg/d level and, to a lesser extent, the 100 and 30 mg/kg/d levels, with the investigations indicating the kidney as the main target organ for toxicity. The microscopic pathol. examination of this organ showed papillary epithelial hyperplasia and/or collecting duct epithelial hyperplasia, with cortical scarring and occasional mineralization in the papilla. Histopathol. changes in the liver, centrilobullar hepatocyte enlargement (accompanied by fine vacuolization) and foci of eosinophilic hepatocytes were considered to reflect the induction of drug-metabolizing enzymes in the liver. Other tissues showing treatment-related findings included the salivary glands, urinary bladder, spleen, pancreas and adrenals. Addnl. other notable findings included (in the high dosage males only) a decline in body weight (from week 34), lower erythrocytic characteristics and slightly higher plasma urea nitrogen and alkaline phosphatase values. The results in this study, therefore, indicated that the non-toxic effect level

L32 ANSWER 37 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:570406 HCAPLUS Full-text

DOCUMENT NUMBER: 121:170406

was 10 mg/kg/d of nefiracetam.

TITLE: Thirteen-week oral toxicity study of the new

cognition-enhancing agent Nefiracetam in dogs

AUTHOR(S): Sugawara, T.; Kato, M.; Suzuki, N.; Akahane,

K.; Takayama, S.

CORPORATE SOURCE: Drug Safety Research Center, Dailchi Pharmaceutical

Co. Ltd., Tokyo, Japan

SOURCE: Arzneimittel-Forschung (1994), 44(2A), 217-19

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE: Journal LANGUAGE: English

The 13-wk oral toxicity of nefiracetam (N-(2,6-dimethylphenyl)-2-(2-oxo-1-pyrrolidinyl) acetamide, DM-9384, CAS 77191-36-7), a new cognition-enhancing agent, was investigated in Beagle dogs of both sexes. No change was observed in the 20 mg/kg group. Hypospermatogenesis and slightly increased deposition of hemosiderin in the spleen were seen in male dogs treated with 60 mg/kg or more. Dosing at 180 mg/kg induced a decrease in food consumption, increases in urinary volume and urinary protein, and renal papillary necrosis in both male and female dogs. The non-toxic dose was 20 mg/kg under these exptl. conditions.

L32 ANSWER 38 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:548981 HCAPLUS Full-text

DOCUMENT NUMBER: 121:148981

TITLE: Thirteen-week oral toxicity study of the new

cognition-enhancing agent nefiracetam in rats

AUTHOR(S): Jindo, T.; Shimizu, Y.; Kato, M.; Takayama,

S.

CORPORATE SOURCE: Drug Safety Research Center, Daiichi Pharmaceutical

Co., Ltd., Tokyo, Japan

SOURCE: Arzneimittel-Forschung (1994), 44(2a), 214-16

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE:

Journal

LANGUAGE: English

Thirteen-week toxicity of nefiracetam (N-(2,6-dimethylphenyl)-2-(2-oxo-1-AB pyrrolidiny1) acetamide, DM-9384, CAS 77191-36-7) was examined in rats by oral administration of 30, 120, or 480 mg/kg. Rats receiving 480 mg/kg showed salivation, prone position, increased water consumption, increased levels of serum total cholesterol, total protein, albumin and total bilirubin, increased liver weight and hypertrophy of liver cells. This hypertrophy of hepatocytes with increased liver weight was also observed in males at 120 mg/kg. The nontoxic dose of nefiracetam under the present exptl. conditions was therefore determined as 30 mg/kg.

L32 ANSWER 39 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1994:595740 HCAPLUS Full-text

DOCUMENT NUMBER:

121:195740

TITLE:

Single dose toxicity study of the new

cognition-enhancing agent nefiracetam in mice, rats,

and dogs

AUTHOR(S):

Sugawara, T.; Kato, M.; Furuhama, K.; Inage,

F.; Suzuki, N.; Takayama, S.

CORPORATE SOURCE:

Drug safety Res. Cent., Daiichi Pharmaceutical Co.,

Ltd., Tokyo, Japan

SOURCE:

Arzneimittel-Forschung (1994), 44(2A), 211-13

CODEN: ARZNAD; ISSN: 0004-4172

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Single oral dose toxicity of nefiracetam (N-(2,6-dimethylphenyl)-2-(2-oxo-1-dimethylphenyl))AB pyrrolidiny1) acetamide, DM-9384, CAS 77191-36-7), a new cognition-enhancing agent, was examined in ddY mice, SD rats, and beagle dogs. The LD530 values of nefiracetam were 2005 mg/kg for male mice and 1940 mg/kg for female mice, 1182 mg/kg for male rats and 1408 mg/kg for female rats, and more than 500 mg/kg for beagle dogs. Common toxic signs in all species were a decrease in locomotor activity, lying on the side or back and loss of righting reflex, considered to be caused by depression of the central nervous system. Pathol., no remarkable change associated with nefiracetam administration was observed in any species. In addition, toxicities of the decomposition product (D-2)and byproducts (Bis, 3-Me and 4-Me) of nefiracetam were examined by oral administration, and of the metabolites (M-3 and M-11) by i.v. injection in male ddY mice. The clin. signs in mice treated with these byproducts were similar to those caused by nefiracetam.

L32 ANSWER 40 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:581898 HCAPLUS Full-text

DOCUMENT NUMBER:

117:181898

TITLE:

Photochromic fulgides

INVENTOR(S):

Tomota, Akihiko; Suzuki, Hisao; Kaneko, Akira

; Tsuboi, Hideki

SOURCE:

Yamaha K. K., Japan

Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT ASSIGNEE(S):

DATE APPLICATION NO. KIND DATE PATENT NO.

JP 04134079 A 19920507 JP 1990-251473 19900920 PRIORITY APPLN. INFO.: JP 1990-251473 19900920

GI

The title compds. I [R1, R3, R5-8 = H, halo, alkyl, alkoxy, aryl, aryloxy, allyl, aralkyl, amino, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, cyano, nitro, ester, CF3; R2, R4 = pyrrolyl, indolyl, NR10R11, Q; X = O, NR9; R9 = H, alkyl, aryl, allyl, aralkyl; R10-11 = H, alkyl, aryl; R12 = alkoxy, amino, (di)alkylamino, (di)arylamino; R13-14 = H, alkyl, aryl; l, n = 0-5] are claimed. I show good light and thermal stability and durability and are useful for optical memory, optical recording, displays, etc.

L32 ANSWER 41 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:540754 HCAPLUS Full-text

DOCUMENT NUMBER:

117:140754

TITLE:

Oxazole fulgide photochromic compound

INVENTOR(S):

Tomota, Akihiko; Suzuki, Hisao; Kaneko, Akira

; Tsuboi, Hideki

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04128282	Α	19920428	JP 1990-247783	19900918
PRIORITY APPLN. INFO.:		•	JP 1990-247783	19900918

OTHER SOURCE(S):

MARPAT 117:140754

GI For diagram(s), see printed CA Issue.

The compound consists of I (R1 = H, halo, alkyl, aryl, allyl, alkoxy, aryloxy, aralkyl, amino, monoalkylamino, dialkylamino, pyrrolyl, indolyl, Q, Q1; R7 = alkyl, alkoxy, amino, monoalkylamino, dialkylamino; l, m = 0-5; R8 = alkyl, aryl, aralkyl, allyl; A = 5- or 6-membered heterocyclic group, etc.; n = 0-4; R2-3 = H, halo, alkyl, alkoxy, aryl, allyl, aryloxy, aralkyl, amino, monoalkylamino, dialkylamino, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, cyano, NO2, ester, trifluoromethyl; X = 0, NR6; R6 = H, alkyl, allyl, aryl, aralkyl; R4R5: = adamantylidene, dicyclopropylmethylidene). The compound showed absorption at long wavelength and is useful for recording using semiconductor laser.

L32 ANSWER 42 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:561036 HCAPLUS Full-text

DOCUMENT NUMBER:

117:161036

TITLE:

Styryl fulgide photochromic compound

Tomota, Akihiko; Suzuki, Hisao; Kaneko, Akira INVENTOR(S):

; Tsuboi, Hideki

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

LANGUAGE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

Δ

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. DATE KIND DATE \_\_\_\_\_\_

JP 04128281 PRIORITY APPLN. INFO .:

PATENT NO.

19920428

JP 1990-247784 JP 1990-247784

19900918 19900918

OTHER SOURCE(S):

MARPAT 117:161036

GT

$$R^{5}n$$
  $X$   $R^{3}$   $R^{4}$   $R^{4}$ 

The composition consists of I (R1-4 = H, halo, alkyl, aryl, allyl, alkoxy, AB aryloxy, allyloxy, aralkyl, amino, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, cyano, NO2, ester, trifluoromethyl; R5 = styryl; n = 1, 2; X = 0, NR6; R6 = H, alkyl, allyl, aryl, aralkyl; Z = O, S, NR6). The compound showed absorption at long wavelength and is useful for recording using semiconductor laser.

HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 43 OF 67

ACCESSION NUMBER:

1992:561035 HCAPLUS Full-text

DOCUMENT NUMBER:

117:161035

TITLE:

Aldehyde intermediates for photochromic compounds Tomota, Akihiko; Suzuki, Hisao; Kaneko, Akira

INVENTOR(S):

; Tsuboi, Hideki

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 04128255	Α	19920428	JP 1990-251472	19900920
PRIORITY APPLN. INFO.:			JP 1990-251472	19900920
GI			·	

$$R^2$$
 $R^1$ 
 $R^6$ 
 $R^6$ 
 $R^7$ 
 $R^7$ 
 $R^7$ 
 $R^7$ 
 $R^7$ 
 $R^7$ 

The intermediate consists of I (R1, R3, R5 = H, alkyl, aryl, allyl, OH, alkoxy, aryloxy, aralkyl, amino, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, cyano, NO2, ester group, trifluoromethyl; R2, R4 = Q; R7 = alkoxy, amino, monoalkylamino, dialkylamino, monoarylamino, diarylamino; l, m = 1-5; R6 = H, alkyl, aryl, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, aralkyl). An obtained fulgide compound from the intermediate showed absorption at long wavelength and is useful for recording using semiconductor lasers.

L32 ANSWER 44 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:633888 HCAPLUS Full-text

DOCUMENT NUMBER:

117:233888

TITLE: .

New building blocks, 3,5-dihydro-1H-thieno[3,4-c]pyrrole 2,2-dioxides; preparation and their

Diels-Alder reaction with dimethyl

acetylenedicarboxylate

AUTHOR(S):

Ando, Kaori; Kankake, Mutuo; Suzuki, Takayoshi;

Takayama, Hiroaki

CORPORATE SOURCE:

Fac. Pharm. Sci., Teikyo Univ., Sagamiko, 199-01,

Janan

SOURCE:

Journal of the Chemical Society, Chemical

Communications (1992), (16), 1100-2

CODEN: JCCCAT; ISSN: 0022-4936

DOCUMENT TYPE:

LANGUAGE:

Journal English

GΙ

V

CO<sub>2</sub>Me

New building blocks, 3,5-dihydro-1H-thieno[3,4-c]pyrrole 2,2-dioxides I (R = PhCH2, Me, PhCH2O2C, 4-MeC6H4SO2, Bz) were prepared and treated with di-Me acetylenedicarboxylate to give new compds.: 7-aza-2,3- dimethylenenorbornene II, the 1:2 adduct III, 1a,3a,6,9- tetrahydrobenz[g]indole IV, and dihydroindolosulfolene V depending on the reaction conditions and the N-substituent.

IT 144425-31-0P 144425-35-4P 144425-37-6P 144425-38-7P 144425-39-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and Diels-Alder reaction of, with acetylenedicarboxylate)

RN 144425-31-0 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(phenylmethyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-35-4 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole-5(3H)-carboxylic acid, phenylmethyl ester, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-37-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-[(4-methylphenyl)sulfonyl]-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-38-7 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 5-benzoyl-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 144425-39-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-methyl-, 2,2-dioxide (9CI) (CA INDEX NAME)

IT 144425-36-5P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 144425-36-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

L32 ANSWER 45 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:72371 HCAPLUS Full-text

DOCUMENT NUMBER:

116:72371

TITLE:

Photochromic fulgide compounds

INVENTOR(S):

Tomota, Akihiko; Kaneko, Akira; Suzuki,

Hisao; Tsuboi, Hideki

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 8 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 03151383	Α	19910627	JP 1989-290579	19891108
PRIORITY APPLN. INFO.:			JP 1989-290579	19891108

GI For diagram(s), see printed CA Issue.

Photochromic fulgides I [Rl = alkyl, alkoxy, NH2, (di)alkylamino, pyrrolyl, indolyl, (CH:CH)lC6H5-mR7m, Q; A = 5- or 6-membered heterocyclic ring or condensed ring containing the heterocyclic ring; R2-5 = H, halo, alkyl, alkoxy, aryl, aryloxy, allyl, aralkyl, amino, pyrrolyl, furyl, thienyl, thiazolyl, oxazolyl, cyano, NO2, ester, CF3; X = O, NR6; R6 = H, alkyl, allyl,

aryl, aralkyl; R7 = alkoxy, NH2, (di)alkylamino; R8 = alkyl, aryl, aralkyl, allyl; l, m = 0-5; n = 0-4] are claimed. I show good thermal stability in the dark and durability and are useful for optical memory, optical recording, displays, etc.,. Thus, acetylacetone was treated with NaNO2 to give (MeCO)2C:NOH, which was treated with PhCHO and the resulting 4-acetyl-5-methyl-2-phenyloxazole (II) N-oxide hydrochloride was stirred with Zn to give II. Treatment of II with di-Et isopropylidenesuccinate, hydrolysis of the resulting half ester, dehydration-cyclization with Ac2O, and photoisomerization gave I (R1 = Ph, R2-5 = Me, X = O) (III). III in PhMe colored with maximum absorption 462 nm by cyclization on irradiation with 365 nm UV and decolored on irradiation with  $\geq 390$  nm visible light.

L32 ANSWER 46 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1990:601448 HCAPLUS Full-text

DOCUMENT NUMBER:

113:201448

TITLE:

Photochromic fulgides containing heterocycles

INVENTOR(S):

Suzuki, Hisao; Kaneko, Akira; Ishizuka,

Mitsuo; Tomota, Akihiko

PATENT ASSIGNEE(S):

Yamaha Corp., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 02160779	A	19900620	JP 1988-314673	19881213
PRIORITY APPLN. INFO.:			JP 1988-314673	19881213
GI				

The title photochromic fulgides I (R = C5-31 alkyl; X = O, S), II, and III are claimed. The above compds. show higher durability and faster response than conventional photochromic fulgides containing furan, thiophen, or *pyrrole* ring instead of oxazole or thiazole in I, pyrazole in II, or indole in III and are useful for optical recording, optical memory, copying materials, etc. Acetone

was acylated with Me(CH2)16CO2Et and the resulting Me(CH2)16COCH2COMe was treated with NaNO2 to give Me(CH2)16COC(:NOH)COMe, which was treated with PhCHO to give 2-phenyl-5-methyl-4-stearoyloxazole. This was treated with EtOCOC(:CMe2)CH2CO2Et and the resulting half ester was hydrolyzed, followed by dehydration to give I (R = heptadecyl, X = 0) (IV) and its (Z)-isomer. IV in toluene colored with maximum absorption 465 nm by cyclization on irradiation with 365 nm UV and decolored on irradiation with  $\geq$ 390 nm visible light and a poly(Me methacrylate) film containing IV showed high durability in repeated use.

L32 ANSWER 47 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:448196 HCAPLUS Full-text

DOCUMENT NUMBER:

111:48196

TITLE:

Photochromic fulgide compounds

INVENTOR(S):

Tomota, Akihiko; Kaneko, Akira; Ishizuka,

PATENT ASSIGNEE(S):

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE ·
JP 01034983	 А	19890206	JP 1987-191025	19870730
JP 06013512	В	19940223		•
PRIORITY APPLN. INFO.:			JP 1987-191025	19870730

MARPAT 111:48196 OTHER SOURCE(S):

For diagram(s), see printed CA Issue. GT

The title compds. having the structure I (R1-R5 = H, halo, alkyl, alkoxy, AB aryloxy, allyl, aralkyl, pyrrolyl, furyl, thienyl, thiazolyl, NH2, CN, NO2, ester, CF3; X = 0, NR8; R8 = H, alkyl, allyl, aryl, aralkyl) are prepared I show excellent thermal stability and are useful in optical recording materials, optical memories, copying, etc. Thus, chlorination of 2,3pentanedione with SO2Cl2 gave 4-chloropentane-2,3-dione, which was refluxed with PhCSNH2 in EtOH to give 4-acetyl-5-methyl-2-phenylthiazole (II). Then, a mixture of II and di-Et isopropylidenesuccinate was added to Me3COK-Me3COH, . then the resulting half ester was hydrolyzed in KOH-EtOH and cyclized by Ac20 to give (Z)-I (R1 = Ph; R2-R5 = Me), 140 mg of which was isomerized into the(E)-isomer by photolysis. A solution of the (E)-isomer in PhMe colored by UV irradiation, returned to its original state by irradiation with light (≥390 nm), and a poly(Me methacrylate) film containing the (E)-isomer showed good thermal stability.

L32 ANSWER 48 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN 1989:448195 HCAPLUS Full-text ACCESSION NUMBER:

DOCUMENT NUMBER:

111:48195

TITLE:

SOURCE:

Photochromic dithiazolylethylene compounds Tomota, Akihiko; Kaneko, Akira; Ishizuka,

INVENTOR(S):

Mitsuo; Suzuki, Hisao

PATENT ASSIGNEE(S):

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

#### PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01034973 PRIORITY APPLN. INFO.:	A	19890206	JP 1987-191026 JP 1987-191026	19870730 19870730
OTHER SOURCE(S):	MARPAT	111:48195		

GI For diagram(s), see printed CA Issue.

AB The title compds. of the structure I (R1-R6 = H, halo, alkyl, alkoxy, aryl, aryloxy, aralkyl, allyl, pyrrolyl, furyl, thienyl, NH2, CN, NO2, thiazolyl, ester, amide, CF3) are prepared I show excellent heat stability and are useful in optical recording materials, optical memories, copying, etc. Thus, refluxing 2,3-pentanedione and SO2Cl2 in CCl4 gave 4-chloropentane-2,3-dione, which was treated with PhCSNH2 in EtOH under reflux to give 4-acetyl-5-methyl-2-phenylthiazole (II). Then, a solution of II in THF was treated with TiCl4 and Zn under reflux for 2 h to give I (R1, R5 = Ph; R2-R4, R6 = Me) (III). A solution of poly(Me methacrylate) and III in cyclohexanone was applied on a glass plate to form a 1.0-μm film, which was colored by UV irradiation, returned to its original state by irradiation with light (≥390 nm), and showed good thermal stability.

L32 ANSWER 49 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1989:448194 HCAPLUS Full-text

DOCUMENT NUMBER:

111:48194

TITLE:

INVENTOR(S):

Photochromic dipyrrolylethylene compounds Tomota, Akihiko; Kaneko, Akira; Ishizuka,

Mitsuo; Suzuki, Hisao

PATENT ASSIGNEE(S):

SOURCE:

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 6 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01034958 PRIORITY APPLN. INFO.:	Α	19890206	JP 1987-191027 JP 1987-191027	19870730 19870730
OTHER SOURCE(S):	MARPAT	111:48194	01 1907 191027	130,0.00
GI .				

The title compds. of the structure I (R1, R3-R7, R9, R10 = H, halo, alkyl, alkoxy, aryl, aryloxy, aralkyl, allyl, pyrrolyl, furyl, thienyl, NH2, CN, NO2, ester, amide, CF3; R2, R8 = CN, NO2, CF3) are prepared I show excellent stability in repeated recording and are useful in optical recording materials, optical memories, copying, etc. Thus, refluxing 5-cyano-1,2-dimethylpyrrole

in Ac2O containing H2SO4 for 1 h gave 3-acetyl-5-cyano-1,2-dimethylpyrrole, which was treated with TiCl4 and Zn under reflux for 2.5 h to give 9.3% cis-1,2-dimethyl-1,2- bis(5-cyano-1,2-dimethyl-3-pyrrolyl) ethene (II). A solution of poly(Me methacrylate) and II in cyclohexanone was applied on a glass plate to form a 1.0- $\mu$ m film, which was colored by UV irradiation and returned to its original state by irradiation with light ( $\geq$ 450 nm).

L32 ANSWER 50 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:448192 HCAPLUS Full-text

DOCUMENT NUMBER:

111:48192

TITLE:

Photochromic fulgide compounds

INVENTOR(S):

Kaneko, Akira; Ishizuka, Mitsuo; Suzuki,

Hisao; Tomota, Akihiko

PATENT ASSIGNEE(S):

SOURCE:

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 01022872 PRIORITY APPLN. INFO.:	A	19890125	JP 1987-177546 JP 1987-177546	19870716 19870716
GI				

$$R^3$$
 $R^2$ 
 $R^3$ 
 $R^4$ 
 $R^5$ 
 $R^5$ 
 $R^5$ 
 $R^6$ 
 $R^7$ 
 $R^7$ 

The title compds. of the structure I (R1, R3-R7 = H, halo, alkyl, aryl, aralkyl, alkoxy, aryloxy, pyrryl, furyl, thienyl, NH2, CN, NO2, CF3; R2 = CN, NO2, CF3, ester, carbamoyl; X = O, NR8; R8 = H, alkyl, aryl, aralkyl) are useful in optical recording materials, optical memories, copying, etc. Thus, a mixture of 3-acetyl-5-cyano-1,2-dimethylpyrrole and di-Et isopropylidenesuccinate was added to a solution of NaH in PhMe, then the resulting half ester was hydrolyzed in KOH-EtOH and cyclized by Ac2O to give (E)-I (R1, R4-R7 = Me; R2 = CN; R3 = H; X = O), whose solution in PhMe was colored upon exposure to UV irradiation and returned to its original state upon irradiation with light (≥ 500 nm).

L32 ANSWER 51 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:604810 HCAPLUS Full-text

DOCUMENT NUMBER:

111:204810

TITLE:

Cadmium sulfide particle doped polymer films for

nonlinear optics

AUTHOR(S):

Ohashi, Y.; Ito, H.; Hayashi, T.; Nitta, A.; Matsuda,

H.; Okada, S.; Nakanishi, H.; Kato, M.

CORPORATE SOURCE:

Cent. Res. Inst., Mitsui Toatsu Chem., Inc., Yokohama,

SOURCE:

Springer Proceedings in Physics (1989), Volume Date

1988, 36(Nonlinear Opt. Org. Semicond.), 81-4

CODEN: SPPPEL; ISSN: 0930-8989

DOCUMENT TYPE:

Journal

English LANGUAGE: AB

CdS particle doped polymer films were prepared by a precipitation-in-gel method in which CdS particles were precipitated in aqueous swollen polymeric gels of crosslinked N-acryloyl-pyrrolidine copolymers by the double decomposition reaction between Cd(OAc)2 and Na2S. Ultrafine particles of average size 30-60 Å were formed in the polymer films. The absorption edge was blue-shifted with the decrease of particle size. Third-harmonic generation (THG) was investigated with 1.06-1.50  $\mu m$  pulsed laser light. Even by the THG measurements, the doped film showed a resonant-enhancement effect.

HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 52 OF 67

ACCESSION NUMBER:

1989:222671 HCAPLUS Full-text

DOCUMENT NUMBER:

110:222671

TITLE:

Flugide photochromic derivative

INVENTOR(S):

Kaneko, Akira; Ishizuka, Mitsuo; Tomota,

Akihiko

PATENT ASSIGNEE(S):

Yamaha Motor Co., Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

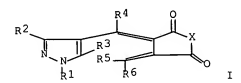
GI

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 63166879	Α `	19880711	JP 1986-313397	19861227
PRIORITY APPLN. INFO.:			JP 1986-313397	19861227
OTHER SOURCE(S):	MARPAT	110:222671		



The title photochromic derivative is I [R1-R6 = H, halo, alkyl, aryl, aralkyl, AΒ alkoxy, aryloxy, pyrroly1, furyl, ethynyl, amino; X = 0, NR7 (R7 = H, alkyl, aryl, aralkyl)]. I [R1 = R2 = R3 = R4 = R5 = R6 = methyl; X = 0] showed  $\lambda max$ = 568 nm at coloration which was higher than that of the conventional photochromic material.

HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 53 OF 67 ACCESSION NUMBER: 1989:222670 HCAPLUS Full-text

DOCUMENT NUMBER:

110:222670

TITLE:

Fulgide photochromic derivative

INVENTOR(S): Kaneko, Akira; Ishizuka, Mitsuo; Tomota,

Akihiko

PATENT ASSIGNEE(S):

SOURCE:

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 9 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 63166878 PRIORITY APPLN. INFO.:	Α	19880711	JP 1986-313400 JP 1986-313400	19861227 19861227

OTHER COHRCE(C).

OTHER SOURCE(S):

MARPAT 110:222670

GΙ

$$R^3$$
 $R^4$ 
 $R^4$ 
 $R^6$ 
 $R^7$ 

The title photochromic derivative is I [R1-R7 = H, halo, alkyl, aryl, aralkyl, alkoxy, aryloxy, pyrrolyl, furyl, ethynyl, amino; X = 0, NR8 (R8 = H, alkyl, aryl, aralkyl) (except for R1 = R4 = R5 = R6 = R7 = Me and X = 0)]. I [R1 = R2 = R4 = R5 = R6 = R7 = methyl; R3 = H; X = 0] showed  $\lambda$ max = 594 nm at coloration which was higher than that of the conventional photochromic material.

L32 ANSWER 54 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:222672 HCAPLUS Full-text

DOCUMENT NUMBER:

110:222672

TITLE:

Fulgide photochromic derivative

INVENTOR(S):

Kaneko, Akira; Ishizuka, Mitsuo; Tomota,

Akihiko

PATENT ASSIGNEE(S):

SOURCE:

Yamaha Motor Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 7 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 63166877 PRIORITY APPLN. INFO.:	A	19880711	JP 1986-313398 JP 1986-313398	19861227 19861227
OTHER SOURCE(S):	MARPAT	110:222672		

$$\begin{array}{c|c}
R^3 & O \\
\hline
R^2 & X \\
R^4 & R^5
\end{array}$$

The title photochromic derivative is I [R1-R5 = H, halo, alkyl, aryl, aralkyl, AB alkoxy, aryloxy, pyrroly1, furyl, ethynyl, amino; X = 0, NR6 (R6 = H, alkyl, aryl, aralkyl)]. I [R1 = R2 = R3 = R4 = R5 = CH3; X = 0] showed  $\lambda max = 585$  nm at coloration which was higher than that of the conventional photochromic material.

L32 ANSWER 55 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:104653 HCAPLUS Full-text

DOCUMENT NUMBER:

110:104653

TITLE:

Photochemical fatigue resistances and thermal

stabilities of heterocyclic fulgides in PMMA film

AUTHOR(S):

Kaneko, Akira; Tomoda, Akihiko; Ishizuka, Mitsuo; Suzuki, Hisao; Matsushima, Ryoka

CORPORATE SOURCE:

Basic Res. Dev. Lab., Yamaha Co., Toyooka, 438-01,

Japan

SOURCE:

Bulletin of the Chemical Society of Japan (1988),

61(10), 3569-73

CODEN: BCSJA8; ISSN: 0009-2673

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Thermal stabilities of the colored forms at 80° and photochem. fatigue AB resistances against coloration-discoloration cycles were measured in PMMA thin films using 14 derivs. of fulgide with different heterocyclic structures. indolyl, oxazolyl, and thiazolyl derivs. showed very low thermal degradabilities (TD ≤1.5%/d) whereas the indolyl and pyrazolyl derivs. showed low photochem. fatigues (PF ≤0.2%/cycle), as compared with the furyl, thienyl, and pyrrolyl derivs. However, many of them featured low photoresponsiveness and low absorptivities for coloration on UV irradiation

HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 56 OF 67

ACCESSION NUMBER:

CORPORATE SOURCE:

1987:458773 HCAPLUS Full-text

DOCUMENT NUMBER:

107:58773

TITLE:

Synthesis of 1-(o-substituted-phenyl)-3,4-

dimethylenepyrrolidines by the thermal elimination of

sulfur dioxide

AUTHOR(S):

Ottenbrite, Raphael M.; Chin, Henry; Alston, Peter V.

Dep. Chem., Virginia Commonw. Univ., Richmond, VA,

23284, USA

SOURCE:

Journal of Heterocyclic Chemistry (1986), 23(6),

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

Journal

LANGUAGE:

English

OTHER SOURCE(S):

CASREACT 107:58773

GI

Precursors of 1-(o-substituted-phenyl)-3,4-dimethylenepyrrolidines I (R = 2-MeOC6H4, 2-MeC6H4, 2-BrC6H4), namely 5-(o-substituted-phenyl)-1,3,4,6-tetrahydrothieno[3,4-c]pyrrole 2,2-dioxides II, were synthesized by reacting RNH2 with 3,4-bis(bromomethyl)-2,5-dihydrothiophene 1,1-dioxide. A disubstitution product along with the desired II was obtained when excess amine was used to neutralize the amine salts that were formed from nucleophilic substitution. A 1,4-HBr elimination product was obtained in three out of four cases when sodium carbonate was used to neutralize the amine salts. The 1,4-HBr elimination product resulted from a competing base attack on the acidic sulfolene protons. The 3,4- dimethylenepyrrolidines were obtained by thermal elimination of sulfur dioxide from II.

IT 109346-47-6P 109346-48-7P 109346-49-8P

RN 109346-47-6 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(2-methoxyphenyl)-, 2,2-dioxide (9CI) (CA INDEX NAME)

RN 109346-48-7 HCAPLUS CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(2-methylphenyl)-, 2,2-dioxide

1H-Thieno[3,4-c]pyrrole, 3,5-dlhydro-5-(2-methylpho (9CI) (CA INDEX NAME)

RN 109346-49-8 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 5-(2-bromophenyl)-3,5-dihydro-, 2,2-dioxide (9CI) (CA INDEX NAME)

L32 ANSWER 57 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1981:551494 HCAPLUS Full-text

DOCUMENT NUMBER:

95:151494

TITLE:

Immobilization of heme complex by radiation

polymerization

AUTHOR(S):

Nishide, H.; Kato, M.; Tsuchida, E.

CORPORATE SOURCE:

Dep. Polym. Chem., Waseda Univ., Tokyo, 160, Japan European Polymer Journal (1981), 17(5), 579-81

SOURCE:

CODEN: EUPJAG; ISSN: 0014-3057

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Transparent films containing Fe-porphyrin complexes were prepared by the  $\gamma$ irradiation of aqueous solns. of a water-soluble monomer (e.g., 1vinylpyrrolidone or 2-hydroxyethyl methacrylate) and complexes of heme, deuteroheme, hemin, or deuterohemin with a 2-methyl-1-vinylimidazole- 1vinylpyrrolidone copolymer. The Fe(II)-porphyrin complex was immobilized in the film by covalent bonding without denaturation, under anaerobic conditions or by protection of the complex with CO. After irradiation, the central Fe ion was reduced spontaneously to the Fe2+ state. Films containing the complex quant. adsorbed CO.

L32 ANSWER 58 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1978:443136 HCAPLUS Full-text

DOCUMENT NUMBER:

89:43136

TITLE:

Optically active N-alkyllactam esters

INVENTOR(S):

Wakabayashi, Toshio; Watanabe, Kenzo; Kato, Yoshinori;

Kato, Masahiko

PATENT ASSIGNEE(S):

Teijin Ltd., Japan

SOURCE:

Jpn. Kokai Tokkyo Koho, 4 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
JP 53025557 JP 61010466	A B	19780309 19860329	JP 1976-98584		19760820
PRIORITY APPLN. INFO.:		19000329	JP 1976-98584	A	19760820

$$\begin{array}{c|cccc}
 & CH_2)_n & & & CH_2CO_2R & & CH_2CO_2H & \\
\hline
 & CH_2CO_2R & & & & CH_2CO_2H & \\
\end{array}$$

Optically active I (n = 1, 2; R = Me, Et) were prepared by treating optically AB active II with RI and NaH in DMF. Thus, 286 mg (+)-II (n = 1) in DMF was treated with 201 mg NaH and then 1.136 g MeI and stirred 7 h at room

temperature to give 294 mg (-)-I (n = 1, R = Me). Also prepared were (-)-I (n = 1, R = Et) and (+)-I (n = 2, R = Me).

L32 ANSWER 59 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1978:443293 HCAPLUS Full-text

ACCESSION NUMBER: 1978:443293 DOCUMENT NUMBER: 89:43293

TITLE: Synthetic approaches to fused heteroaromatic compounds

by the condensation reactions of functional pyrroles

AUTHOR(S): Uchida, Takane

CORPORATE SOURCE: Fac. Educ., Fukui Univ., Fukui, Japan

SOURCE: Journal of Heterocyclic Chemistry (1978), 15(2), 241-8

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 89:43293

GΙ

Diacyl- and triacylpyrroles, obtained by one pot synthesis from aziridines and acetylenic dipolarophiles, underwent condensation reactions. On treatment of 3,4-di- and 2,3,4-tribenzoylpyrroles with hydrazine hydrate and phosphorus pentasulfide, pyrrolopyridazine derivs., e.g. I, and fused thiophenes, e.g. II, resp., were prepared The structure proofs for I were based on the 13C FT-NMR spectrum of the corresponding 13C-enriched compds.

IT 66864-55-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 66864-55-9 HCAPLUS

CN 7,11-Epithio-7H-pyrrolo[3',4':5,6]isoindolo[2,1-a]quinoxaline-8,10(7aH,9H)-dione, 10a,11-dihydro-12-(4-nitrophenyl)-6,7,9,11-tetraphenyl- (9CI) (CA INDEX NAME)

ACCESSION NUMBER: 1977:422129 HCAPLUS Full-text

DOCUMENT NUMBER: 87:22129

TITLE: On the aromatic stability of positional isomers

consisting of bicyclic systems composed entirely of

five-membered heterocycles

AUTHOR(S): Milun, M.; Trinajstic, N.

CORPORATE SOURCE: Rudjer Boskovic Inst., Zagreb, Yugoslavia SOURCE: Croatica Chemica Acta (1977), 49(1), 107-13

CODEN: CCACAA; ISSN: 0011-1643

DOCUMENT TYPE: Journal LANGUAGE: English

AB Bicyclic conjugated compds. consisting entirely of 5-membered heterocyclic rings are studied by topol. resonance energy (TRE) index. The TRE correctly

predicts the aromatic behavior of the positional isomers.

IT 63156-10-5

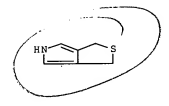
RL: PRP (Properties)

(aromaticity of, topological resonance energy and graph theory in

relation to)

RN 63156-10-5 HCAPLUS

CN 1H-Thieno[3,4-c]pyrrole, 3,5-dihydro- (9CI) (CA INDEX NAME)



L32 ANSWER 61 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1974:505350 HCAPLUS Full-text

DOCUMENT NUMBER: 81:105350

TITLE: Synthesis of nonclassical thiophenes

AUTHOR(S): Potts, K. T.; McKeough, D.

CORPORATE SOURCE: Dep. Chem., Rensselaer Polytech. Inst., Troy, NY, USA

SOURCE: Journal of the American Chemical Society (1974),

96(13), 4268-75

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB P2S5 treatment of suitable vicinal dibenzoyl heterocycles was established as a convenient pathway to three nonclassical thiophene systems: tetraphenylthieno[3,4-c]thiophene (I), 5-methyl-1,3,-4,6-tetraphenylthieno[3,4-c]pyrrole (II), as well as hexaphenylthieno[3,4-f]isothionaphthene (III). 1,3-Dipolar cycloaddn. reactions with PhCOC.tplbond.CCOPh, utilizing the "masked" 1,3-dipole of several mesoionic systems, readily provided the precursors to the  $10\pi$ -electron heterocycles. II formed 1:1 primary cycloadducts with activated olefins across both the 4 and 6 positions (azomethine ylide) and 1 and 3 positions (thiocarbonyl ylide), these addns. being examples of kinetic and thermodynamic product control. In some

cases, the thiocarbonyl ylide adducts underwent thermal elimination of the elements of H2S giving rise to bicyclic heteroaromatics. The addition of PhCOC.tplbond.CCOPh occurred only across the azomethine ylide affording a stable 1:1 adduct. I and PhCOC.tplbond.CCOPh formed an unstable 1:1 adduct which decomposed by the elimination of S forming 5,6-dibenzoyl-1,3,4,7-tetraphenylisothionaphthene which, in turn, afforded III upon treatment with P4S10. This novel  $14\pi$ -electron system underwent cycloaddn. reactions with olefins across the 1 and 3 positions.

41223-66-9P 52579-55-2P IT

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

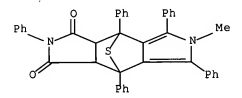
41223-66-9 HCAPLUS RN

4,7-Epithio-2H-isoindole-5,6-dicarbonitrile, 4,5,6,7-tetrahydro-2-methyl-CN 1,3,4,7-tetraphenyl-,  $(4\alpha,5\alpha,6\beta,7\alpha)$ - (9CI) (CA INDEX NAME)

Relative stereochemistry.

52579-55-2 HCAPLUS RN

4,8-Epithiobenzo[1,2-c:4,5-c']dipyrrole-1,3(2H,3aH)-dione, CN 4,6,8,8a-tetrahydro-6-methyl-2,4,5,7,8-pentaphenyl-,  $(3a\alpha, 4\alpha, 8\alpha, 8a\alpha)$  - (9CI) (CA INDEX NAME)



HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 62 OF 67

ACCESSION NUMBER: 1974:133307 HCAPLUS Full-text

DOCUMENT NUMBER: 80:133307

Nonclassical condensed thiophenes. IV. Derivatives TITLE:

of thieno[3,4-c]furan-SIV and thieno[3,4-c]pyrrole-SIV Cava, Michael P.; Sprecker, Mark A.; Hall, William Roy

AUTHOR(S):

Dep. Chem., Univ. Pennsylvania, Philadelphia, PA, USA CORPORATE SOURCE:

Journal of the American Chemical Society (1974), SOURCE:

96(6), 1817-21

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE: Journal LANGUAGE: English

AB

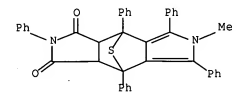
For diagram(s), see printed CA Issue. GI

Tetraphenylthieno-[3,4-c]furan-SIV (I) has been generated as a transient intermediate which could be trapped efficiently by dimethyl acetylenedicarboxylate. In contrast, the red N-methyl-1,3,4,6-tetraphenylthieno[3,4-c]pyrrole (II) and pentaphenylthieno[3,4-c]-pyrrole have been synthesized and are stable in the solid state. Some simple reactions of the thienopyrrole system are discussed including catalytic reduction oxidation and cycloaddns. The results of CNDO/2 calcns. for the parent heterocycles thieno-[3,4,-c]thiophene-SIV, thieno[3,4-c]furan-SIV, and thieno[3,4-c]-pyrrole-SIV are presented and correlated with the observed chemical of these systems.

IT 52579-55-2P

RN 52579-55-2 HCAPLUS

CN 4,8-Epithiobenzo[1,2-c:4,5-c']dipyrrole-1,3(2H,3aH)-dione, 4,6,8,8a-tetrahydro-6-methyl-2,4,5,7,8-pentaphenyl-,  $(3a\alpha,4\alpha,8\alpha,8a\alpha)$ - (9CI) (CA INDEX NAME)



L32 ANSWER 63 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1973:159489 HCAPLUS Full-text

DOCUMENT NUMBER:

78:159489

TITLE:

Thieno[3,4-c]pyrrole system, a tetravalent sulfur

heterocycle showing coth azomethine ylide and thiocarbonyl ylide dipolar characteristics

AUTHOR(S):

Potts, K. T.; McKeough, D.

CORPORATE SOURCE:

Dep. Chem., Rensselaer Polytech. Inst., Troy, NY, USA

SOURCE:

Journal of the American Chemical Society (1973),

95(8), 2749-50

CODEN: JACSAT; ISSN: 0002-7863

DOCUMENT TYPE:

LANGUAGE:

Journal English

GI For diagram(s), see printed CA Issue.

The thienopyrrole I was prepared by cyclizing BzMeNCHPhCO2H with PhCOC.tplbond.CCOPh to give 3,4-dibenzoyl-1-methyl-2,5-diphenylpyrrole and treating this with P2S5, followed by 10% NaOH. Treatment of I with fumaronitrile gave the 1:1-cycloadduct II in refluxing PhMe and III in refluxing C6H6. II eliminated H2S on heating to give 1,3,4,7-tetraphenyl-5,6-dicyano-2-methylisoindole.

IT 41223-66-9P 41688-99-7P

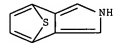
RN 41223-66-9 HCAPLUS

CN 4,7-Epithio-2H-isoindole-5,6-dicarbonitrile, 4,5,6,7-tetrahydro-2-methyl-1,3,4,7-tetraphenyl-,  $(4\alpha,5\alpha,6\beta,7\alpha)$ - (9CI) (CA INDEX NAME)

Relative stereochemistry.

41688-99-7 HCAPLUS RN

CN 4,7-Epithio-2H-isoindole (9CI) (CA INDEX NAME)



HCAPLUS COPYRIGHT 2006 ACS on STN L32 ANSWER 64 OF 67

ACCESSION NUMBER:

1974:3407 HCAPLUS Full-text

DOCUMENT NUMBER:

80:3407

TITLE:

Synthesis of 1-aryl-3,4-dimethylenepyrrolidines by the

thermal elimination of sulfur dioxide

AUTHOR(S):

Ottenbrite, Raphael M.; Alston, Peter V.

CORPORATE SOURCE:

Dep. Chem. Pharm. Chem., Virginia Commonw. Univ.,

Richmond, VA, USA

SOURCE:

Journal of Heterocyclic Chemistry (1973), 10(5),

785-90

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

LANGUAGE:

Journal English

CASREACT 80:3407

OTHER SOURCE(S):

For diagram(s), see printed CA Issue.

Reaction of 3,4-bis(bromomethyl)-2,5-dihydrothiophene 1,1-dioxide (I) with AΒ arylamines gave 5-aryl-1,3,4,6-tetrahydrothieno[3,4-c]pyrrole 2,2-dioxides (II) in good yields for arylamines with substituent groups with Hammett  $\sigma$ values <0.40. II were thermally decomposed to the corresponding 1-aryl-3,4dimethylenepyrrolidines (III) in good yields.

50872-65-6P 50872-66-7P IT

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

50872-65-6 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-(4-methylphenyl)-, 2,2-dioxide CN (CA INDEX NAME)

RN 50872-66-7 HCAPLUS

1H-Thieno[3,4-c]pyrrole, 3,5-dihydro-5-phenyl-, 2,2-dioxide (9CI) (CA CN INDEX NAME)

L32 ANSWER 65 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN 1972:72340 HCAPLUS Full-text ACCESSION NUMBER:

76:72340 DOCUMENT NUMBER:

1-(p-Chlorophenyl)-3,4-dimethylenepyrrolidine, a new TITLE:

pyrrole isomer

Gschwend, Heinz W.; Haider, Hasan AUTHOR(S):

CIBA Pharm. Co. Div., CIBA-GEIGY Corp., Summit, NJ, CORPORATE SOURCE:

Journal of Organic Chemistry (1972), 37(1), 59-61 SOURCE:

CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE:

Journal English

LANGUAGE:

CASREACT 76:72340

OTHER SOURCE(S): For diagram(s), see printed CA Issue.

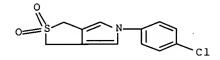
1-(p-Chlorophenyl)-3,4-dimethylenepyrrolidine (I), isomeric with the AB corresponding pyrrole, was prepared The key step involves the SO2 extrusion of 5-(p-chlorophenyl)-1,3,4,6-tetrahydrothieno [3,4-c]-pyrrole 2,2-dioxide (II) under reduced pressure. The structure of the diene was confirmed by its spectral data as well as by conversion to a dimer and two Diels-Alder adducts.

32515-68-7P IT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

32515-68-7 HCAPLUS RN

1H-Thieno[3,4-c]pyrrole, 5-(4-chlorophenyl)-3,5-dihydro-, 2,2-dioxide CN (9CI) (CA INDEX NAME)



L32 ANSWER 66 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER:

1968:418890 HCAPLUS Full-text

DOCUMENT NUMBER:

69:18890

TITLE:

Synthetic studies of benzocycloheptene derivatives. Syntheses and stereochemical consideration of

trimethoxybenzocycloheptenone derivatives Hayashi, Yuji; Kato, Masahiko; Miwa, Toshio;

Sakan, Takeo

CORPORATE SOURCE:

Osaka City Univ., Osaka, Japan

SOURCE:

Nippon Kagaku Zasshi (1967), 88(5), 569-73

CODEN: NPKZAZ; ISSN: 0369-5387

DOCUMENT TYPE:

Journal

LANGUAGE:

AUTHOR(S):

Japanese

GI For diagram(s), see printed CA Issue.

 $trans-\beta-[2-(3,4,5-Trimethoxyphenyl)]$  ethyl]paraconic acid lactone (I) (1.0 g.) AB and 18 g. HF was heated in an autoclave 13 hrs. at 50  $\pm$  5° to give 442 mg. trans-6,7,8,9-tetrahydro-2,3, 4-trimethoxy-7-hydroxy-5-oxo-5Hbenzocyclohepten-6-ylacetic acid lactone (trans-II), m. 157-8°, and 13 mg. 8,9-dihydro- 2,3,4-trimethoxy-5- oxo- 5H-benzocyclohepten-6-ylacetic acid (III). Similarly, cis-I gave 37.1% cis-II, m. 157-8°. Heating trans-II with

HCl in EtOH gave cis-II. trans-II (435 mg.) in 13 ml. EtOH was treated with NaOEt prepared from 33 mg. Na and 1.5 ml. EtOH to give 404 mg. III, m. 183°. Similarly, cis-II gave III. Heating III with 2N HCl and EtOH yielded cis-II. III was treated with CH2N2 to give the Me ester, m. 121-2°, which (606 mg.) was heated with 20 ml. EtOH saturated with NH3 at 50° to yield 479 mg. cis-7-amino-6,7,8,9-tetrahydro-2, 3,4-trimethoxy-5-oxo-5H- benzocyclohepten-6-ylacetic acid lactam (cis-IV), m. 186-7°. cis-IV was treated with 15% aqueous HBr in AcOH 30 hrs. at room temperature to give 45.5% cis-7-amino-6,7,8,9-tetrahydro-2,3-dimethoxy-4- hydroxy-5-oxo-5H-benzocyclohepten-6-ylacetic acid lactam (cis-V), m. 236°. Similar treatment of cis-IV with HBr at 50-5° yielded 50% trans-V, m. 242°; acetate m. 217-19°. Attempted Reformatskii reaction of cis-IV yielded trans-IV, m. 200-200.5°, and cis-V and 39% cis-IV were recovered. cis-V isomerized partially to trans-V on heating. Methylations of trans-V and cis-V with CH2N2 were followed by uv spectrography. trans-V reacted .apprx.50% in 70 hrs. to give cis-IV, whereas cis-V reacted completely in 15 hrs. to give cis-IV.

L32 ANSWER 67 OF 67 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER:

1967:443173 HCAPLUS Full-text

DOCOMENT

67:43173

TITLE:

Photolysis of 2,3-diphenyl-2-cyclopropenylcarboxylic

acid azide and its homolog

AUTHOR(S):

Castellucci, N. C.; Kato, Masahiko; Zenda,

Hiroshi; Masamune, Satoru

CORPORATE SOURCE:

Univ. Alberta, Edmonton, Can.

SOURCE:

Chemical Communications (London) (1967), (10), 473-4

CODEN: CCOMA8; ISSN: 0009-241X

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GI For diagram(s), see printed CA Issue.

During photolysis, I converted to α-ketocarbene and II underwent an intramol. addition to the double bond. In a further study of the addition reaction using III and IV 0.2% in ether at -30°, photolysis of III yielded 50-60% isocyanate (V), 4% diphenylacetylene, and 15% amide (VI), m. 162-3. V was also obtained by a normal Curtius rearrangement of III at 70°. Similarly, photolysis of IV, followed by treatment with EtOH at 70°, gave 6% 2,5-diphenylpyrrole (VII), m. 140.5-1.5, 5% 5,6-diphenylpyridone (VIII), m. 272-3°, 1% diphenylacetylene, 15% corresponding urethane (IX), and 21% 3,4-diphenylpyridone (X). Protonation occurred readily between EtOH and the isocyanate (XI) obtained thermally from IV. The resulting cation reacted with the solvent, or the double bond of the cyclopropene to form IX, and X. The photolysis of XI gave VII. XII was considered an intermediate in the formation of VII and possibly VIII.

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(FILE 'HOME' ENTERED AT 14:38:37 ON 18 DEC 2006)

FILE 'REGISTRY' ENTERED AT 14:38:43 ON 18 DEC 2006

L\*\*\* DEL STRUCTURE UPLOADED
L1 STRUCTURE UPLOADED

L2 0 SEA SSS SAM L1

D QUE L1

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L4
             1 SEA ABB=ON PLU=ON (L3 OR L4)
L5
               D SCAN
               SEL RN L5
     FILE 'REGISTRY' ENTERED AT 14:40:18 ON 18 DEC 2006
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L6
               152940-73-3/BI OR 22722-98-1/BI OR 2526-64-9/BI OR 63156-10-5/B
                I OR 646065-36-3/BI)
               D SCAN
            79 SEA SSS FUL L1
L7
               SAVE L7 DEBORAHO50/A TEMP
              2 SEA ABB=ON PLU=ON L7 AND L6
L8
              5 SEA ABB=ON PLU=ON L6 NOT L8
L9
                D SCAN
     FILE 'HCAPLUS' ENTERED AT 14:41:45 ON 18 DEC 2006
            2 SEA ABB=ON PLU=ON L8
L10
            32 SEA ABB=ON PLU=ON L7
L11
            32 SEA ABB=ON PLU=ON (L10 OR L11)
L12
            31 SEA ABB=ON PLU=ON L12 NOT L5
L13
     FILE 'MEDLINE, EMBASE, BIOSIS, CAOLD, WPIX' ENTERED AT 14:42:22 ON 18 DEC
     2006
             O SEA ABB=ON PLU=ON L7
L14
     FILE 'MEDLINE, EMBASE, BIOSIS, CAOLD' ENTERED AT 14:42:40 ON 18 DEC 2006
          O SEA ABB=ON PLU=ON L7
L15
     FILE 'HCAPLUS' ENTERED AT 14:42:57 ON 18 DEC 2006
                E KATO M/AU
           1265 SEA ABB=ON PLU=ON ("KATO M"/AU OR "KATO M A M F"/AU OR "KATO
                M J"/AU OR "KATO M K"/AU OR "KATO M T"/AU OR "KATO M TAKAYUKI"/
                AU OR "KATO MASAHIKO"/AU)
               E KANEKO A/AU
          301 SEA ABB=ON PLU=ON ("KANEKO A"/AU OR "KANEKO AKIRA"/AU)
L17
           4 SEA ABB=ON PLU=ON L16 AND L17
L18
            4 SEA ABB=ON PLU=ON (L18 OR L5)
L19
            34 SEA ABB=ON PLU=ON (L16 OR L17) AND ?PYRROL?
L20
               D KWIC
             36 SEA ABB=ON PLU=ON (L18 OR L19 OR L20)
L21
             36 SEA ABB=ON PLU=ON L21 NOT L13
L22
     FILE 'MEDLINE, EMBASE, BIOSIS, CAOLD, WPIX' ENTERED AT 14:45:01 ON 18 DEC
     2006
          23488 SEA ABB=ON PLU=ON KATO M?/AU
L23
          2736 SEA ABB=ON PLU=ON KANEKO A?/AU
L24
           78 SEA ABB=ON PLU=ON L23 AND L24
2 SEA ABB=ON PLU=ON L25 AND PYRROL?
179 SEA ABB=ON PLU=ON (L23 OR L24) AND PYRROL?
L25
L26
L27
            56 SEA ABB=ON PLU=ON (L23 OR L24) AND (PYRROL?(L) DERIVATIVE#)
L28
            51 DUP REM L28 (5 DUPLICATES REMOVED)
L29
                    ANSWERS '1-4' FROM FILE MEDLINE
                     ANSWERS '5-11' FROM FILE EMBASE
                    ANSWERS '12-51' FROM FILE WPIX
            51 SEA ABB=ON PLU=ON (L26 OR L29)
L30
```

FILE 'STNGUIDE' ENTERED AT 14:46:58 ON 18 DEC 2006

D QUE L22

D QUE L30

D QUE L13

FILE 'HCAPLUS, MEDLINE, EMBASE, WPIX' ENTERED AT 14:47:17 ON 18 DEC 2006 L31 110 DUP REM L22 L30 L13 (8 DUPLICATES REMOVED)

ANSWERS '1-67' FROM FILE HCAPLUS ANSWERS '68-71' FROM FILE MEDLINE ANSWERS '72-78' FROM FILE EMBASE ANSWERS '79-110' FROM FILE WPIX

D OUE L26

D QUE L22

D QUE L13

67 DUP REM L26 L22 L13 (2 DUPLICATES REMOVED)

ANSWERS '1-2' FROM FILE WPIX

ANSWERS '3-67' FROM FILE HCAPLUS

D ALL ABEQ TECH L32 1-2

D IBIB ABS HITSTR RETABLE L32 3-67

FILE HOME

L32

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 15 DEC 2006 HIGHEST RN 915749-75-6 DICTIONARY FILE UPDATES: 15 DEC 2006 HIGHEST RN 915749-75-6

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH June 30, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

### FILE HCAPLUS

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FILE COVERS 1907 - 18 Dec 2006 VOL 145 ISS 26 FILE LAST UPDATED: 17 Dec 2006 (20061217/ED)

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This file contains CAS Registry Numbers for easy and accurate substance identification.

#### FILE MEDLINE

FILE LAST UPDATED: 15 Dec 2006 (20061215/UP). FILE COVERS 1950 TO DATE.

All regular MEDLINE updates from November 15 to December 16 have been added to MEDLINE, along with 2007 Medical Subject Headings (MeSH(R)) and 2007 tree numbers.

The annual reload will be available in early 2007.

This file contains CAS Registry Numbers for easy and accurate substance identification.

#### FILE EMBASE

FILE COVERS 1974 TO 18 Dec 2006 (20061218/ED)

EMBASE is now updated daily. SDI frequency remains weekly (default) and biweekly.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BIOSIS FILE COVERS 1969 TO DATE. CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1969 TO DATE.

RECORDS LAST ADDED: 14 December 2006 (20061214/ED)

FILE CAOLD FILE COVERS 1907-1966

FILE LAST UPDATED: 01 May 1997 (19970501/UP)

This file contains CAS Registry Numbers for easy and accurate substance identification. Title keywords, authors, patent assignees, and patent information, e.g., patent numbers, are now searchable from 1907-1966. TIFF images of CA abstracts printed between 1907-1966 are available in the PAGE display formats.

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file supports REGISTRY for direct browsing and searching of all substance data from the REGISTRY file. Enter HELP FIRST for more information.

FILE WPIX

FILE LAST UPDATED: 13 DEC 2006 <20061213/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200680 <200680/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

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FILE STNGUIDE

FILE CONTAINS CURRENT INFORMATION.

LAST RELOADED: Dec 8, 2006 (20061208/UP).